

Methods of genetically modifying donor cells by gene transfer for grafting into the central nervous system to treat diseases of damaged cells are described. The modified donor cells produce a molecule capable of affecting the recovery of cells in the CNS.

(57) Abstract

(54) Title: GRAFTING GENETICALLY MODIFIED CELLS TO TREAT DISORDERS OF THE CENTRAL NERVOUS SYS.

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Attempts to repair the mammalian brain or replace CNS functions resulting from defects or following disease damage to the CNS are hampered by an incomplete understanding of the complex structure-function relationships between cells of the CNS. Although knowledge of some basic principles in the CNS, such as their interactions between different types of cells or systems and cellular circuitries in different regions of the brain and their relationship to the outside world, have been gained, far behind these approaches are the large number of regions of the brain and their relationships to the outside world.

Background of the invention

The present invention relates to the use of recombinant technology for genetic modification of donor cells for grafting into the central nervous system (CNS) of a subject to treat defects, disease or damage of the CNS. More specifically, the invention relates to the insertion of a gene encoding a molecule having ameliorative effects on cells including neurons into donor cells such that when the donor cells are grafted into the CNS, they express the molecule into damaged cells.

Technical field of the invention

This invention was made with government support under Grant Contract No. NIA-06088 awarded by the Office of Naval Research, and Grant Contract Nos. HD-20034, NS-24279, HD-00669 awarded by NIH. The Government has certain rights in this invention.

Acknowledgment of Government Support

GRAFTING GENETICALLY MODIFIED CELLS TO TREAT DISEASES OF THE CENTRAL NERVOUS SYSTEM

different cell types in the mammalian CNS and the number and complexity of their connections. In addition, the blood-brain barrier makes access to the brain for diagnosis, treatment and the design of new therapies more difficult.

In spite of the absence of sophisticated knowledge of pathophysiology of most normal or abnormal brain functions, some attempts at pharmacological therapy for CNS dysfunctions have already become useful and effective. These include the use of psycho-active drugs for psychiatric disorders such as schizophrenia, and specific repackaging in the rare cases in which the bio-chemical and cellular bases of the CNS disorder are relatively better understood, as in Parkinson's disease. At the core of most therapeutic approaches is the objective replacement or repairing a specific chemical function of the brain that has been lost as a consequence of disease or damage.

In spite of the absence of sophisticated knowledge of pathophysiology of most normal or abnormal brain functions, some attempts at pharmacological therapy for CNS disorders include the use of psycho-active drugs for psychiatric disorders such as schizophrenia, and specific replacement therapy in the rare cases in which the biochemical and cellular bases of the CNS disease are relatively better understood, as in Parkinson's disease. At the core of most therapeutic approaches is the objective of replacing or reacting off-reactivating a specific chemical function of the brain that has been lost as a consequence of dis-

Intracerebral neural grafting has emerged recently as an additional potential approach to CNS therapy. The replacement or addition of cells to the CNS are able to produce and secrete therapeutically useful metabolites may offer the advantage of avoiding repeated drug administration which also avoids avoiding the drug delivery system. While the (Rosenthalin, Science 235:772-774 (1987)). While the concept is basic procedures of intracerebral grafting have been known for decades, most of the factors that optimize the survival of grafted cells have only recently come to be investigated and partially understood. (Björklund et al., in Neural Grafting in the Mammalian CNS, p. 709, Elsevier, Amsterdam (1985); Sladek et al., in Neural Transplants: Development and Function, Plenum Press, NY (1984)). Several factors critical for reliable

Parkinson's disease is an age-related disorder characterized by a loss of dopamine neurons in the substantia nigra of the midbrain, which have the basal ganglia as their major target organ. The symptoms include tremor, rigidity and ataxia. The disease is progressive but can be treated by replacement of dopamine through the administration of pharmacological doses of L-dopa.

Level of experimental clinical application in Parkinson's disease. This approach has reached a therapeutic level of CNS disease. For clinicians for the design of function and potentiality for neurologists in the study of CNS reliable tool for neurologists to come a valid and optimized, intracerebral grafting has become a recognized and As these critical factors have become recognized and

(6) Vasculization: it is critical that the grafts be vascularized rapidly or otherwise sufficiency well nourished from the environment.

(5) The importance of target-donor matching: neurons survive better when they are grafted to a site which they would normally innervate.

(4) Immunological response: the brain is not totally an immunologically privileged site.

(3) Availability of neurotrophic factors in the host and donor tissue: wound-induced neurotrophic factors enhance graft survival.

(2) Age of the host: young recipients accept grafts more readily than older ones.

(1) Age of the donor: efficiency of grafting is reduced with increasing age of donor cells.

including the following, and effective graft survival have been identified,

(1) the chemical deficit is well known (dopamine),
plantation of genetically engineered cells, because
Parkinson's disease is a candidate disease for the trans-
With this background, it seems likely that

the vicinity of the defective cells.
cells constitutively producing and secreting dopamine in
functional graft and that it may be sufficient to have
that synaptic connectivity may not be a requisite for a
Neurosci., 6:266-270 (1983)). These experiments suggest
Acad. Sci., 457:53-81 (1986); Dunnett et al., Trends
dopaminergic neurotoxins. (Bjorklund et al., Ann. N.Y.
reversing the behavioral deficits induced by selective
in Parkinson's disease, has been shown to be effective in
cell bodies and also the area of the brain most affected
nigra, an area of the brain rich in dopamine-containing
cells such as fetal brain cells from the substantia
316:831-836 (1987)). The transplantation of other donor
62:169-173 (1985); Madrazo et al., New Eng. J. Med.
affected patients. (Bjorklund et al., J. Neurosurg.
homografting adrenal medullary cells to the brain of
diseased basal ganglia of Parkinson's patients by
provide the neurotransmitter dopamine to cells of the
been used in humans. Several attempts have been made to
brain for which therapeutic intracerebral grafting has
Parkinson's disease is the first disease of the

from the precursor.
the remaining cells cannot synthesize sufficient dopamine
that the patients reach a threshold of cell loss, wherein
development of the refractory state, but the simplest is
L-DOPA. There are many suggested mechanisms for the
often become refractory to the continued effect of
although with chronic use of pharmacotherapy the patients
Clinical Neurology P. 185, Elsevier, Amsterdam (1986),
Neurosci. 9:512 (1986); Vincken et al., in Handbook of
the precursor for dopamine, L-DOPA, (Marsden, Trends

The recent demonstration of genetic components in a rapidly growing list of other CNS diseases, including Huntington's disease, (Gusella et al., *Nature* 306:234-238 (1983)) Alzheimer's disease, (DeLaBar et al., *Science*, N.Y. 235:1390-1392 (1987); Goldgaber et al., *Science*, N.Y. 235:877-880 (1987); St. George-Hyslop et al., *Science*, N.Y. 235:885-890 (1987); Tanzi et al., *Science*, N.Y. 235:880-884 (1987)); bipolar disease (Baron et al., *Nature* 326:289-292 (1987)); schizophrenia (Sherrington et al., *Nature* 336:164-167 (1988) and many other major human diseases, suggests that these other CNS diseases will eventually become accessible to gene therapy approaches.

(2) the human and rat genes for the rate-limiting enzyme in the production of dopamine have been cloned (tyrosine hydroxylase), (3) the anatomical localization of the enzyme in the sympathetic neurons has been determined (synaptosomal fraction), and (4) synaptic connectivity does not appear to be required for complete functional restoration.

A great deal of attention has recently been paid to the use of gene delivery vectors derived from murine retroviruses (Andersson, *Science* 226:401-409 (1984); Gilboa et al., *Biotecniques* 4:504-512 (1986)) for gene transfer at later stages of development. Gene transfer in vitro using such retroviral vectors is extremely efficient for a broad range of recipient cells, the vectors have a suitable capacity for added genes, and infection with them does little metabolic damage to recipient cells. (Shimotohno et al., Cetl 26:67-77 (1981); Wei et al., J. Virol. 39:935-944 (1981); Tabain et al., Molec. Cell. Biol. 2:426-436 (1982)). Several useful systems have demonstrated that the expression of genes introduced into cells by means of retroviral vectors can correct metabolic aberrations in vitro in several human genetic diseases associated with single-gene enzyme deficiencies. (Ranchoff et al., *Proc. Natl. Acad. Sci. USA* 83:6563-6567 (1986); Wilkins et al., *J. Biol. Chem.* 259:7842-7849 (1984)). There has been particular interest in bone marrow as a potential target organ for this approach to gene therapy because of the prevalence and importance of disorders of bone marrow-derived cell lineage in a variety of major human diseases, including thalassemias and sickle-cell anemia, Gaucher's disease, chronic granulomatous disease (CGD) and immunodeficiencies resulting from deficiencies of the purine pathway enzymes, adenosine deaminase (ADA) and purine nucleoside phosphorylase (PNP) (ADA)

and imunodeficiencies resulting from deficiencies of target organs, such as the liver, have also recently become theoretical for genetic manipulation through the demonstration of infection of cells from such organs with viral vectors (Moltef et al., *Proc. Natl. Acad. Sci. USA* 84:3344-3348 (1987)). Furthermore, the

There are several ways to introduce a new function into target cells in the CNS in a phenotypically useful way i.e. to treat defects, disease or dysfunction (Fig. 1). The most direct approach, which bypasses the need for cellular grafting entirely, is the introduction

A recent study developed model of gene therapy uses target cells removed from a subject, placed in culture, genetically modified in vitro, and then re-implemented into the same subject (Mollett et al., Rheumatologic Dis. Chin. N. Amer. 14(2):459-477 (1988); Egliitis et al. Biotechniques 6:608-614 (1988); Ledley, J. Pediatr. 110:1-8 (1987)). Target cells have incited bone marrow stem cells (Joyner et al., Nature 305:556-558 (1983); Milller et al., Science 225:630-632 (1984); Williams et al., Nature 310:476-480 (1984)); fibroblasts (Seldeen et al., Science 236:714-718 (1982)); Graver et al., Proc. Nat'l. Acad. Sci. USA 84:1050-1054 (1987) and St. Louis et al., Proc. Nat'l. Acad. Sci. USA 85:3150-3154 (1988)), keratinocytes (Morgan et al., Science 237:1476-1479 (1987)) and hepatocytes (Mollett et al., Proc. Nat'l. Acad. Sci. USA 84:3344-3348 (1987)). This indirect approach of in vivo gene transfer is necessitated by the inability to transfer genes effectively directly into cells in vivo. Although there has been some recent progress towards gene delivery using modified neurons in culture (Gellet et al., Science 241:1667-1669 (1988)), this indirect approach of gene delivery using neurons in culture (Gellet et al., in vivo gene transfer has not yet been applied to the gene delivery system (GNS). CNTS.

disscovery of numerous cell-specific regulatory signals such as *cis*-acting enhancers, tissue-specific promoters and other sequences may provide tissue specific gene expression in many other organs even after general specific infections and gene transfer in vivo (Khoury et al., Cell 33:313-314 (1983); Serflin et al., Trends Genet. 1:224-230 (1985)).

There are several types of neurons in the mammalian brain. Cholinergic neurons are found within the mammalian brain. Cholinergic neurons are found within the mammalian brain and project from the medial septum and

Fig. 1).
cells to provide a functional new transgene (5, in progeny viruses that might in turn infect nearby target tent helper viruses, could produce locally high titers of replicative vector but also replication-complementary, an introduced donor cell infected with not only finally, a final product USA 83:9231-9235 (1986)).
al., Proc. Nat'l. Acad. Sci. USA 83:9231-9235 (1986).
damage (Hefte, J. Neurosci. 6:2155 (1986); Williams et protection of cholinergic neurons after follow-on CNS postulated with CNS cells, as in the case of NGF-mediated damage (Fig. 1). This type of "co-operative" has been demonstrated with CNS cells, as in the case of NGF-mediated alternative they may be genetically modified *in vivo* (4, the donor cells may be genetically modified *in vitro* or and used by nearby defective target cells (3, in Fig. 1).
secrete a diffusible gene product that can be taken up although it has not yet been demonstrated conclusively in neurons. Still other donor cells could express and has been called "metabolic co-operation" and is known to occur between fibroblasts and glial cells (Gruener et al., Proc. Nat'l. Acad. Sci. USA 82:6662-6666 (1985)).
Biology. Acta. 560:1-66 (1979)). This process permits the efficient diffusion of metabolically important small molecules from one cell to another, leading to phenotypic changes in the recipient cell (Lowenstein), although a genetic modification donor cell that could function is expressed in defective target cells by introducing a new Parkinson's disease (1, in Fig. 1). Alternatively, a new Tay-Sach's disease, possibly Lesch-Nyhan disease, and Parkin's disease (1, in Fig. 1). Alternative, a new function is aberrant as a consequence of a developmental or genetic defect, i.e. normal cells in the case of of a transgene directly into the cells in which that

It would be advantageous to develop procedures for gene transfer via efficient vectors followed by

Studies have shown that chronic intra-ventricular administration of NGF before axotomy will prevent collateral ingrowth of neurons in the septum (Heftei, 1986). Gage et al. (1987) and Williams et al. (1986) reported that the same treatment regimen can also prevent retrograde degeneration of motoneurons in the lumbar spinal cord after peripheral nerve lesion. In contrast, Kromer and Schiene (1986) found no evidence of retrograde degeneration in motoneurons after dorsal root section. These findings indicate that the retrograde degeneration of motoneurons after peripheral nerve lesion is not a general phenomenon, but rather depends on the type of lesion and the time interval between the lesion and the onset of regeneration.

vertebral lumb of the diagonal band of Broca to the hippocampal formation in the basal forebrain. The short, nerve-like portion of the brain connecting the medial septum and vertical limb of the diagonal band with the hippocampal formation is termed the "fimbria fornix". The fimbria fornix contains the axons of the neurons of section or lesion (also termed "axotomy"). Axotomy severely affects the cholinergic neurons in the septum and diagonal band and results in the death of up to one-half of the cholinergic neurons (gagé et al., Neuroscience 19:241-256 (1986)). This degenerative response is attributed to the loss of trophic support from nerve growth factor (NGF), which is normally transposed retrogradely in the intact brain from the hippocampus to the septal cholinergic cell bodies (Korsching et al., Proc. Nat'l. Acad. Sci. USA 80:3513 (1983); Whittmore et al., Proc. Nat'l. Acad. Sci. USA 83:817 (1986); Sheltor et al., Proc. Nat'l. Acad. Res. 83:525 (1987); and Seijler et al., Brain Res. 300:33 (1984)).

transcriptase).

(GAG=group specific antigen; Env=envelope; P0L=reverse transmissible retrovirus vectors containing a transgene.

Fig. 3 is a diagrammatic depiction of the preparation of

using genetically modified donor cells.

introducing a new function into target cells in the CNS

Fig. 2 is a diagrammatic representation of strategies for

into target cells.

introducing and analyzing the effect of a new function

Fig. 1 is a diagrammatic representation of methods for

Brief Description of the Drawings

Liorative interactions of injured neurons.

implanting of material to facilitate reconnection or ameliorating disease or damaged cells in the central nervous system. The methods include grafting accompanied by

treatment disease or damaged cells in the central nervous system may be co-administered with a therapeutic agent for injection in suspension into the central nervous system

foreign DNA into a cell. The cells may be cultured and affects the cells, or by other methods of introducing transgene encoding a product which directly or indirectly targets cellular vectors containing an inserted therapeutic effect on the . The cells may be modified using viral or that directly or indirectly provides an ameliorative into the central nervous system to produce a molecule that disrupts or damage of cells in the central nervous system

ous system by grafting genetically modified donor cells

defects, disease or damage of cells in the central nervous system provides methods for treating

Summary of the Invention

in vivo to treat disorders of the CNS.

intracerebral grafting of the genetically modified cells

Fig. 4 is a diagrammatic representation of the linear restriction map of the integrated vectors LSAPALM and PLNH2 as described in Example I, intra (arrows indicate the location of the promoter and the direction of transcription). The location of the hexapeptide added by in vitro mutagenesis, LTR=long terminal repeat).

Fig. 5 is a depiction of the circular restriction map of vector PLNH2 as described in Example I, intra. Fig. 6 is photomicrographs of primary rat fibroblasts previously infected with hypoxanthine quanine phosphoribosyl transferase (HPRT) that have been implanted in rat basal ganglia as described in Example I, intra. (A, a=anti-fibronectin; B, b=creseyl violet; C, c=GFP; magnification: A-C=88X; a-c=440X).

Fig. 7 is photographs of isoelectric focusing gels for Hprt enzymatic activity of brain extracts from basal ganglia as described in Example I, intra.

Fig. 8 is a depiction of the circular restriction map of vector PLLNTL as described in Example I, intra. Fig. 9 is a depiction of the circular restriction map of vector PPR1 as described in Example I, intra. Fig. 10 is a depiction of the circular restriction map of vector PUCRK as described in Example I, intra.

Fig. 11 is a diagrammatic depiction of the linear restriction map of the integrated NGF retroviral vector PLN-BRN1 containing the 777 base pair Hgal-PstI fragment of mouse nerve growth factor (NGF) cDNA under control of the viral long terminal repeat as described in Example II, intra (arrow indicates insertion site; LTR=long terminal repeat; psi (φ) =retroviral example II, intra (arrows indicate transcription initiation).

Fig. 16 is a diagrammatic depiction of the linear restriction map of PLTHRN_L retroviral integrated vector described in Example III, Intra (arrows indicate the location of the promoters and the gene marker).

for A, C and E; magnification: A,B = 20X; C-F = 220X).
B,D,F animal grafted with control cells as described
in of a through the medial septum: E = high power magnification
of a through the medial septum: E = high power magnification
NGF-infected donor cells; C = higher power magnification
power magnification of an animal grafted with
histochemistry as described in Example II, Intra (A = low
Fig. 15 is photomicrographs of acetylcholinesterase

Intra.
ence and absence of NGF as described in Example II,
immunoreactive cells in the septum of a rat in the presence
-
Fig. 14 is a graph showing survival of CHAT-

magnification: A and B = 20X; C and D = 70X, E and F = 220X).
cells; B,D,F = animal with graft of control cells; magnifi-
cation: A,C,E = animal with graft of retrovirus-infected
CHAT; A,C,E = animal with graft of retrovirus-infected
taken through the medial septum of tissue stained for
into the fibrilla fornia cavity; C-F = coronal sections
Intra (A,B = fibronectin staining in fibroblasts grafted
in for fibronectin and CHAT as described in Example II,
Fig. 13 is photomicrographs of immunohistochemical stain-

Intra.
vector PLN.BRN_L as shown in Figure 11 and described in
Fig. 12 is a depiction of the circular restriction map of

neo⁺= neomycin-resistance gene marker.
Packing signal; RSV = Rous sarcoma virus promoter;

Fig. 19 is a graph showing the average percent change in the number of rotations from baseline to post-transplantation in 4 experimental groups of animals as described in Example III, infra (magnification: A,C = 10X; B,D = 20X).
In Example III, infra (magnification: A,C = 10X; B,D = 20X).
Caudate showing fibronectin immunoreactivity as described in Example III, infra (magnification: A,C = 10X; B,D = 20X).

Fig. 18 is photomicrographs of fibroblast grafts to the caudate showing fibronectin immunoreactivity as described in Example III, infra (magnification: A,C = 10X; B,D = 20X).

Fig. 17 is a depiction of the derivation of vector PLTHNL as shown in Fig. 16 and described in Example III, infra.

The strategy for transferring genes into donor cells in vitro is outlined in Fig. 2 and includes the following basic steps: (1) selection of appropriate model "reporter" genes or transgenes whose expression is corrected with CNS disease or dysfunction; (2) selection and development of suitable and efficient vectors for gene transfer; (3) preparation of donor cells from primary cultures or from established cell lines; (4) demonstration that the donor implanted cells expressing the new

Gene Transfer Into Donor Cells In Vitro

The present invention relates to a process for grafting genetic material modified by introduction of a retroviral vector containing a transgene, for example a gene encoding nerve growth factor (NGF) protein. The gene encodes a transgene, for example a central nervous system system, for example, to treat damage sustained by the brain, to treat CNS disease or damage of cells in the CNS, donor cells such as fibroblasts are modified by introduction of a disease or trauma. Preferably, for treating defects, to repair damage sustained by the cells from CNS to modify donor cells to produce a molecule that is capable of directly or indirectly affecting cells in the CNS use of vectors carrying foreign gene inserts (transgenes) the CNS. More particularly, the invention relates to the tral nervous system (CNS) to treat disease or trauma of grafting genetic material modified donor cells into the central nervous system relating to a process for

In order that the invention herein described may be more fully understood, the following detailed description is set forth.

Detailed Description of the Invention

Although other vectors may be used, preferred vectors for use in the methods of the present invention are viral (including retroviral) vectors. The viral vector selected should meet the following criteria: 1) the vector must be able to infect the donor cells and thus viral vectors having an appropriate host range must be selected; 2) the transferred gene should be capable of persisting and being expressed in a cell for an extended period of time without causing cell death for stable maintenance and any damage to target cells. Murine retroviral vectors offer an efficient, useful, and presently the best-characterized means of introducing and expressing foreign genes efficiently in mammalian cells.

Choice of vector

Most of the techniques used to transform cells, consisting of vectors and promoters, are widely practiced in the field, and most practitioners are familiar with the basic procedures. However, for convenience, the following and procedures. However, for convenience, the following

The methods described below to modify donor cells using retroviral vectors and grafting into the CNS are merely for purposes of illustration and are typical of those that might be used. However, other procedures may also be employed, as is understood in the art.

Genetic Modification of Donor Cells

function, are viable and can express the transgene product stably and efficiently; (5) demonstration that the trans-plantation causes no serious deleterious effects; and (6) demonstration of a desired phenotypic effect in the host animal.

(1980).

separations is found in Methods in Enzymology 65:499-560 using standard techniques. A general description of size by polyacrylamide gel or agarose gel electrophoresis is size separation of the cleaved fragments may be performed extractions by precipitation with ethanol. If desired, extraction, and the nucleic acid recovered from aqueous with phenol/chloroform, and may be followed by ether after each incubation, protein is removed by extraction at 37°C are workable, although variations can be tolerated. Incubation times of about one hour to two hours at about used to insure complete digestion of the DNA substrate. solution. Typically, an excess of restriction enzyme is cleaved by one unit of enzyme in about 20 μl of buffer (log.). In general, about 1 μg of plasmid or DNA sequences with the suitable restriction enzyme (or enzymes) under conditions which are generally understood in the art, and the particulates of which are specified by the manufacturer of these commercial avilable restriction turer of these commercial available restriction enzymes.

Site-specific DNA cleavage is performed by treating

are cleaved, tailored, and religated in the form desired. plasmids, DNA sequences, or synthesized oligonucleotides plasmids, DNA sequences, or synthesized oligonucleotides Spring Harbor Laboratory, New York (1982)). Isolated which are well understood in the art (see Materials and Methods Standard ligation and restriction techniques employed the therapeutic gene coding and control sequences desired the construction of suitable vectors containing the

General Methods for Vector Construction

obviously damage their host cells. effectively, and under most conditions do not kill or random sites in the host genome, express genes stably and integrate by reasonably well understood mechanisms into

In vector construction employing "vector fragments", the vector fragment is commonly treated with bacterial alkaline phosphatase (BAP) or calf intestinal alkaline phosphatase (CIP) in order to remove the 5' phosphate and prevent religation of the vector. Digesions are conducted at pH 8 in approximately 150 mM Tris, in the presence of Na⁺ and Mg²⁺ using about 1 unit of BAP or CIP per

total ends concentration.

10-30 fold molar excess of linkers) are performed at 1 μM intermolecular blunt end ligations (usually employing a concentrations (5-100 nM total end concentration).

Ligation are usually performed at 33-100 ng/ml total DNA (for "blunt end" ligation). Intermolecular "sticky end" 1 mM ATP, 0.3-0.6 (Wells) units T4 DNA ligase at 14°C 10 mM-50 mM NaCl, and either 40 μM DTT, 33 mg/ml BSA, Tris-Cl pH 7.5, 10 mM MgCl₂, 10 mM ATP, 0.01-0.02 (Wells) the following standard conditions and temperatures: 20 mM ligations are performed in 15-50 μl volumes under

singlē-stranded portion.

with S1 nucleic acid or Bal-31 results in hydrolysis of any nucleic acid. Treatment under appropriate conditions not precipitated. The mixture is extracted with phenol/chloroform and ethanol precipitation is extracted with phenol/klenow, nature of the sticky ends. After treatment with klenow, selected DNTPs, within the limitations dictated by the formed by supplying only one of the DNTPs, or with are present. If desired, selective repair can be performed by singlē strands, even though the four DNTPs protruding 3', singlē strands, even though the four DNTPs about 15 to 25 min at 20°C to 25°C in 50 mM Tris (pH 7.6) 50 mM NaCl, 6 mM MgCl₂, 6 mM DTT and 5-10 μM DNTPs. The cleotide triphosphates (DNTPs) using incubation times of merase I (Klenow) in the presence of the four deoxyuridine triphosphate (dNTPs) may be blunt ended by treating with the large fragment of E. coli DNA poly-

mg of vector at 60°C for about one hour. In order to recover the nucleic acid fragments, the preparation is extracted with phenol/chloroform and ethanol precipitated. Alternatively, religation can be prevented in vectors which have been double digested by additonal restriction enzymes digestsion of the unwanted fragments.

In a preferred viral vector the transgene is brought under the control of either the viral LTR promoter or of an internal promoter, and retained enhancer signals or of an internal promoter, and retained signals within the retroviral LTR can still bring about efficient integration of the vector into the host cell genome. To prepare transmissible virus (Figure 3), recombinant DNA molecules of such defective vectors are transfected into "producer" cell lines that contain a provirus expressing all of the retroviral functions required for packaging of viral transcripts into transmissible viruses particles, but lacking the crucial packaging signal for encapsidation of RNA transcripts of the provirus into mature virus particles. These include the group specific antigen (GAG) and envelope (ENV) genes which encode capsid proteins and reverse transcriptase (POL). Because of this deletion, transcripts from the producer cannot be packaged into viral particles and the helper (POL). However, an integrated defective retroviral vector introduced into the same cell by means of calcium-activated channels, therefore, generate only empty virus particles.

B10L. 5:431-437 (1985); and Miller, et al., Mol. Cell. Biotechchniques 6:608-614 (1988)) and are now in common use in many laboratories. Retroviral vectors contain long terminal repeats (LTRs) and packaging sequences (psi) sequences, as well as plasmid sequences for replicon insertion in eukaryotic cells. Much of the rest of the viral genome is removed and replaced with other promoters and genes. Vectors are packaged as RNA in viruses particles following cell lines. These include psi (y)2 which produce viral particles that can infect rodent cells and ham and virtual particles that can infect human cells that can affect a broad range of species.

Since herpes viruses are capable of establishing a latent infection and an apparently non-pathogenic relationship with some neural cells, herpes based vectors, e.g. HSV-1, may be used. Since it should be possible to take advantage of an eventual improved understanding of other paramyxoviruses and the human immunodeficiency virus. In most cases, with the exception of retrovirus (HIV), to develop useful delivery and expression vectors. In most cases, with the exception of the CNS effectively, such as rabies virus, measles, mumps, rubella, varicella-zoster, and other paramyxoviruses and the human immunodeficiency virus. These viruses are most pronounced in cells of the nervous system (GAG, ENV and POU genes have been replaced by the transgene (X) with intact psi sequence, produces transcripts that can be packaged in trans since they do contain the packaging sequence. The cells contain 2 proviruses sequences integrated into different sites of the host cell genome. Because RNA transcripts from the newly introduced proviruses contain the packaging sequence they are efficient encapsidated into virus particles by means of viral functions produced in trans. Ideally, the result is the production by the cells of infectious particles part of the infection and possibly probably since it may lead to spreading of other helper virus. In most, but not necessarily all models of gene therapy, the production of helper virus is wild-type helper virus. In most, but not necessarily all models of gene therapy, the production of helper virus is probably undetectable since it may lead to spreading or other tissue in the host animal.

Or other tissue in the host animal. Since it is possible to produce probably undetectable in lymphoid infection and possibly probably since it may lead to spreading of other helper virus. In most, but not necessarily all models of gene therapy, the production of helper virus is wild-type helper virus. In most, but not necessarily all models of gene therapy, the production of helper virus is probably undetectable since it may lead to spreading or other tissue in the host animal.

Because phosphatase-mediated transformation (Graham and Vandendorp, Virology 52:456-467 (1973)) in which the GAG, ENV and POU genes have been replaced by the transgene (X) with the intact psi sequence, produces transcripts that can be packaged in trans since they do contain the packaging sequence. The cells contain 2 proviruses sequences integrated into different sites of the host cell genome. Because RNA transcripts from the newly introduced proviruses contain the packaging sequence they are efficient encapsidated into virus particles by means of viral functions produced in trans. Ideally, the result is the production by the cells of infectious particles part of the infection and possibly probably since it may lead to spreading or other tissue in the host animal.

in the art.

Liposomal mediated transfection and other methods known
transfection (Graham et al., Virol. 52:456-467 (1973));
mediated transfection such as calcium phosphate
et al., Mol. Cell. Biol. 6:703-706 (1986)); chemically
for example, microinjection; electroporation (Toneguzzo
include nonviral physical transfection of DNA into cells;
methods may be used. For example, non-vector methods
inserting foreign DNA transgenes into donor cells other
In addition to the above-described methods for

expression of the vector.

sequences not needed for the infection, stabilization or
reduced through the elimination of all viral regulatory
other cellular sequences. This possibility can be
type virus by recombination with endogenous viruses-like or
or "rescue" of pathogenic, replication-competent, wild-
viral vectors is the potential for the eventual emergence
A possible problem posed by the use of defective

rabies and poliovirus and other human and animal viruses.
e.g. bovine papilloma virus type I (BPV); vaccinia;
viruses; Epstein-Barr virus (EBV); papilloma viruses,
(MMLV); papovaviruses such as JC, SV40, polyoma, adeno-
retroviruses such as Maloney murine leukemia virus
fer into cells for correction of CNS disorders include
Other virus vectors that may be used for gene trans-

and other possible CNS specific functions in the mouse.
protein and glial fibrillary acidic protein (GFAP) genes,
JC virus, glial-specific expression of the proteolipid
that regulate the oligodendroglial-specific expression of
enhancer, promoter and other sequences, such as those
be conferred by the use of appropriate cell-specific

The choice of donor cells for implantation depends heavily on the nature of the expressed gene, characteristics of the vector and DNA synthesis for efficient infection, interaction and gene expression (Weiss et al., eds., Cold Spring Harbor viruses, 2nd Ed., Weiss et al., RNA Tumour viruses, New York (1985)), if such vectors are used the donor cells are probably actively growing cells such as primary fibroblasts culture or established cell lines, donor cells in embryonic neural cells or replicating adult neurons and possibly in selected areas such as the olfactory mucosa and keratinocytes, hepatocytes, connective tissue cells, keratinocytes, hepatocytes, connective tissue suitable donor cells include fibroblasts, neurons, glial cells, epidermal cells, chromaffin cells and other mammalian cells, primary cultures of adult rat hepatocytes, ordinary fibroblasts, vaccinia, or other viruses, as well as the use of efficient, non-viral methods for transduction DNA into donor cells such as the recent developed retroviral vector (Toneguzzo et al.). In addition, the development of many other kinds of vectors derived from herpes, vaccinia, or other viruses, as well as the use of efficient, non-viral methods for transfer into many other cells presently not susceptible to retroviral infection.

Choice of donor cells

Grafting

Mechanisms of Phenotypic Correction by Donor Cells

The methods of the invention contemplate intra-cerebral grafting of donor cells containing a transgene inserted to the region of the CNS having sustained, deficit, disease or trauma.

Neural transplantation or "grafting" involves transplantation of cells into the ventricular cavity of a host brain. Conditions or subdually onto the surface of a host brain. Grafts at the site of transplantation; 2) retention of the graft at the site of transplantation; and 3) minimum amount of pathological reaction at the site of transplantation.

Plantsation include: 1) viability of the implant; 2) retention of the graft at the site of transplantation; and 3) minimum amount of pathological reaction at the site of transplantation.

example embryonic brain tissue, into host brains have been described in *Neural Grafting in the Mammalian CNS*, Bjorklund and Stenovi, eds., (1985) Das, Ch. 3 pp. 23-30; Freed, Ch. 4, pp. 31-40; Stenovi et al., Ch. 5, pp. 41-50; Brundin et al., Ch. 6, pp. 51-60; David et al., Ch. 7, pp. 61-70; Seiger, Ch. 8, pp. 71-77 (1985), incorporated by reference herein. These procedures include intraparenchymal transplantation, i.e. within the host brain (as compared to outside the brain or extraparenchymal transplantation) achieved by injection or deposition (Das, supra).

The two main procedures for intraparenchymal transplantation (Das, supra) are: 1) injecting the donor cells within the host brain parenchyma or 2) preparing a cavity by surgery cal means to expose the host brain parenchyma and then depositing the graft into the cavity (Das, supra). Both

and host brain tissue at the time of grafting, and both facilitate anatomic integration between the graft and host brain tissue. This is of importance if it is required that the graft become an integral part of the host brain and to survive for the life of the host.

Alternatively, the graft may be placed in a ventricle, e.g. a cerebral ventricle or subdurally, i.e. on the surface of the host brain where it is separated from the host brain parenchyma by the intervening pia mater or arachnoid and pia mater. Grafting to the ventricle may be accomplished by injection of the donor cells or by growing the cells in a substrate such as 30% collagen to form a plug of solid tissue which may then be implanted into the ventricle to prevent dislocation of the graft.

For subdural grafting, the cells may be injected around

Injections into selected regions of the host brain may be made by drilling a hole and piercing the dura to permit the needle of a microsyringe to be inserted. The microsyringe is preferably mounted in a stereotaxic frame and three dimensional stereotaxic coordinates are selected for placing the needle into the desired location of the brain or spinal cord.

The cellular suspension procedure thus permits grafting of genetically modified donor cells to any predetermined site in the brain or spinal cord, is relatively non-traumatic, allows multipe grafting simultaneously in several different sites using the same cell

the donor cells must be properly prepared for grafting. For example, for injection of genetic ally modified donor cells according to the present invention, cells such as fibroblasts obtained from skin samples are placed in a suitable culture medium for growth and maintenance of the cells, for example a solution containing fetal calf serum and allowed to grow to confluence. The cells are loosened from the culture substrate for example using a buffered solution such as phosphate buffered saline (PBS) containing 0.05% trypsin and placed in a buffered solution such as PBS supplemented with 1 mg/ml of glu- cose; 0.1 mg/ml of MGCL2; 0.1 mg/ml CACL2 (complete PBS) plus 5% serum to inactivate trypsin. The cells may be

Preparation of Donor Cells

graft in the host brain to prevent isolation or cavity in the host brain to traumatic brain. In addition, the donor cells should possess sufficient growth potential to fill any lesion or cavity in the host brain to prevent isolation or cavity in the host brain to traumatic brain. In addition, the donor cells should be cleaned and bleeded before attempting to graft. In addition, the site of injury must be cleaned and bleeded stopping before grafting different procedures, for example, the site will require different procedures, for example, the site of injury must be cleaned and bleeded before grafting of donor cells into a traumatized brain.

For transplantation into spinal cord grafting, tissue is removed from a transplanted cavity, for example as described by Stenveriet al., super, by removing bone overlying the brain and stopping bleeding with a material such as gelfoam. Suction may be used to create the cavity. The plant may be placed in the cavity. More than one transplant is then placed in the cavity. The cells or solid tissue implants.

suspension, and permits mixtures of cells from different anatomical regions.

The most effective mode and timing of grafting of the transgenic donor cells of the invention to treat defects, disease or trauma in the CNS of a patient will depend on the severity of the defect and on the security and course of disease or injury to cells such as neurons in the CNS, the patient's health and response to

cells during a phase of immune tolerance of the host animal, as in fetal life. It is imperative to minimize the potential for rejection and graft-versus-host reaction by the graft and can be demonstrated to foreign antigens in the rat brain. Recent work has shown conclusively that immune responses can be demonstrated to foreign antigens in the rat brain, but considered to be an immunologically privileged organ, but gene product. The mammalian brain has traditionally been considered to be foreign cells or to the foreign host animal to the host brain to the foreign cells or to the adequate vascularization, or the establishment of mechanisms of cell implantation, on the long-term survival of implanted cells may depend on effects of the viral infection on the cells, on cellular damage produced by the culture conditions, on the mechanics of cell implantation, or the establishment of adquate vascularization, or the immune response of the host brain to the foreign cells.

In addition, the host must be appropriately prepared for grafting of donor cells. This depends on the site of the host brain for grafting.

Washed with PBS using centrifugation and are then resuspended in the complete PBS without trypsin and at a selected density for injection. In addition to PBS, any osmotically balanced solution which is physiologically compatible with the host subject may be used to suspend and inject the donor cells into the host.

clude the development of specific neural connections to
the use of non-neuronal cells for grafting may pre-

infection.
gyrus. Such cells may be suitable targets for retroviral
such as those in the olfactory mucosa and in the dentate
hippocampus, or continue to divide through adulthood,
such as the ventral leaf of the dentate gyrus of the
there are cells within the CNS that are late to develop,
cells after grafting into the brain. Alternatively,
be able to establish synaptic connections with other
able to retain other neural characteristics, they may
retroviral or other viral vectors, and if they are also
might be susceptible to efficient transduction by
et al., Science 241:1667-1669 (1988)). Such neurons
gene transfer and then for in vivo implantation. (Geller
culture systems may soon become available for in vitro
neuronal cells, suggest that replicating neural cells
involving the immortalization of embryonic hippocampal
infection. However, recent studies, including those
refractory to non-replicating neuronal cells to viral
carrying non-transformed cell-culture systems and the
developed and exploited because of the paucity of repli-
tions to study these possibilities have not yet been
fateful intercellular synaptic connections. Model sys-
tems may require the establishment of re-establishment of
dents may important issues of appropriate or fateful gene
the important issues of gene-transfer systems,
Of course, as in all other gene-transfer systems,

cell.
not typical effect and not so high as to be toxic to the
gene expression is sufficient to achieve the desired phe-
expression must be resolved to ensure that the level of
the important issues of appropriate or fateful gene
Of course, as in all other gene-transfer systems,

sional.
treatment and the judgment of the treating health profes-

intracerebrally into a subject to treat disease or
tors carrying a therapeutic transgene are grafted
ferred embodies in which donor cells containing vec-
the methods of the invention are exemplified by pre-

treat effects, disease and injury of the CNS.
genetically modifying donor cells for grafting CNS to
the present invention therefore provides methods for

homogenates of neurons or placenta.
from synthetic or biological materials, for example
or may be used in conjunction with neural bridges formed
axonal regeneration and reconnection of injured neurons,
donor cells may serve as neural bridges to facilitate
Brain Res. Bull. 15:13-18 (1985)). Thus, the grafted
implants of peripheral homogenates of neurons (Wendt,
campus of the brain has also been demonstrated using
(1981)). Connectivity between the septum and hippocampus
of an injured rat (David and Aguayo, Science 214:931-933
join the medulla oblongata and upper thoracic spinal cord
peripheral nerve segments have been used to successfully
Annals of the N.Y. Acad. of Sciences, 495:1-9 (1987)).
Gillford press, pp. 457-484 (1985) and Aguayo et al.,
Aguayo, in *Synaptic Plasticity*, Cotman, ed., New York,
CNA tissues. Neural bridges have been described (see
which facilitate reconnection between neurons in damaged
Alternatively, "neural bridges" may be provided

lizing and clearing a neurotoxin.
toxin "sink" by expressing a new gene product and metabo-
ucts or metabolites. The donor cell may also act as a
uptake by target cells of secreted donor cell gene prod-
tigous junctions ("metabolic co-operation") or through
sion of a required gene product or metabolite, through
cells or target cells *in vivo* would be through the diffu-
typic effects of fibroblast or other non-neuronal donor
resident target cells of the host. Therefore, the pheno-

The methods of the invention also contemplate the use of grafting of transgenic donor cells in combination with other therapeutic procedures to treat disease or trauma in the CNS. Thus, genetically modified donor cells of the invention may be co-grafted with other cells of the CNS, such as chromaffin cells from the adrenal gland, fibroblasts which exert beneficial effects on cells in the CNS, both genetically modified and non-genetically modified cells, both geneticaly modified and non-geneticaly modified cells of the invention may be co-grafted with other cells of the CNS.

In a third preferred embodiment fibroblasts were infected with a retroviral vector, and the modified gene was secreted to express and secrete L-DOPA by fibroblasts. Fibroblasts were grafted into the caudate of rats model-ing Parkinson's disease as a result of unilateral dopamine depletion. The cells survived and produced sufficient L-DOPA to decrease the rotational movement caused by dopamine depletion.

In a second preferred embodiment fibroblasts were implanted into the brains of rats with surgically lesions of the fibrilla coronary region. The grafted cells survived and produced sufficient NGF to prevent the degeneration of cholinergic neurons that would die without treatment. In addition, the protected cholinergic cells sprouted axons that projected in the direction of the cellular source of NGF.

Hprt-deficient rat fibroblast line 208F, primary rat fibroblasts, and postnatal day-1 primary rat astrocytes were used to demonstrate that cultured cells genetically modified using retroviral vectors can survive when implanted in the mammalian brain and can continue to express foreign gene products.

In order that the invention described herein may be more fully understood, the following examples are set forth. It should be understood that these examples are for illustrative purposes only and are not to be construed as limiting the scope of this invention in any manner.

Moreover, the genetic liability modified donor cells of the invention may be co-administered with therapeutic agents useful in treating defects, trauma or diseases of the CNS, such as growth factors, e.g. nerve growth factor; gangliosides; antibiotics; antidiabetics, neurotransmitters, neurotransmitters, toxins, neurite promoting molecules; and anti-metabolites and precursors of these molecules such as the precursor of dopamine, L-DOPA.

geriatrically modified donor cells may thus serve to support the survival and function of the co-grafted, non-geriatrically modified cells, for example fibroblasts modified to produce nerve growth factor (NGF) *in vivo* as described in the Examples, infra.

Donor hypoxanthine guanine phosphoribosyl transferase (Hprt)-deficient 208F rat fibroblast cells (Jolly et al., Proc. Nat'l Acad. Sci., USA 80:477-481 (1983)) were infected with the prototype Hprt vector 225:630-632 (1984); Yee et al., Gene 53:97-104 (1987) PLASPALM expressing Hprt DNA (Miller et al., Science 230:1395-1398 (1985); Wolff et al., Nat'l Acad. Sci. (USA) expressing both the transposon neomycin-resistance gene (neo^r) and the luciferase gene (luc^r) with the Hprt DNA derived from vector PLP2 (Miller et al., Mol. Cell. Biol. 6:2895-2902 (1986)) and containing a novel C-terminal hexapeptide encoding a protein with a novel C-terminal hexapeptide added by in vitro mutagenesis of the translational termination codon (Yee et al., Gene 53:97-104 (1987)). Vector PLASPALM was constructed as follows: Vector PLP2 (Yee et al., Gene 53:97-104 (1987)) was digested with XbaI to yield a 1.3 kb fragment which was then ligated into plasmid PLP2 (Miller, supra) which had also been restricted using XbaI and BamHI. The resulting vector was PLASPALM.

The vector contained the neo^r gene (luc^r gene) and the Hprt gene (luc^r gene) under the control of the firefly luciferase promoter and enhancer of the human gene (neo^r) and the T5 neomycin-resistance gene (Hprt). The 5' and 3' LTRs were derived from the early gene (Hcmv). The 5' and cytosine-galactosidase genes were derived from murine leukemia virus (MLV) as described by Mason et al., in Science 234:1372-1378 (1986)). Vector PLNHL2 was constructed as firefly luciferase (luc^r) and the T5 neomycin-resistance gene (neo^r) and the Hprt gene (luc^r gene) under the control of the Hprt promoter and enhancer of the human gene (neo^r) and the T5 neomycin-resistance gene (Hprt). The vector contained the Hprt gene (luc^r gene) under the control of the firefly luciferase promoter and enhancer of the human gene (neo^r) and the T5 neomycin-resistance gene (Hprt).

Infection of Cells

Intracerebral grafting of genetically modified cells expressing Hprt transgene to the brain

Example I

excised, and examined histologically and biochemically. Areas containing the implanted cells were identified, and after 1 week to 3 months the animals were killed and injected at a rate of 1 μ l/min for a total volume of 3-5 μ l. Between 10,000 and 100,000 cells per microliter were regions of the rat brain using a sterile microsyringe. Saline solution and injection stereotactically into several regions of the rat brain using a balanced glucose-saline solution were resuspended in a balanced glucose-

Grafting

in the rat brain. Serum, to reduce the likelihood of immunological response overnight with serum-free medium or medium containing rat G418-resistant cells were harvested following incubation with the neo^r-luciferase vector only. HAT-resistant and with the neo^r-luciferase vector only. HAT-resistant and primary fibroblasts and astrocytes were infected primarily expressing Hprt and with the neomycin analog G418 for hypoxanthine, amphotericin and thymidine (HAT) for cells expressing neor, respectively, to ensure that only cells expressing neor were used.

The cells were grown in selective medium containing infected cells were used. Cells expressing neor, respectively, to ensure that only cells expressing neor were used.

PLNH2 (Fig. 5). The BamHI restricted together forming vector restricted PSV2A were ligated together forming SSP1 as above. The BamHI restricted PLNH2 and HindIII-SSP1 isolate the luciferase fragment. The ends were repaired San Diego, CA) was restricted with HindIII and SSP1 to PSV2A (dewet et al., Moléc. And Cell Biol., supra, and supplied by) (Dr. Subramani, University of California, the ends were repaired using Klenow polymerase. Plasmid restricted with BamHI to remove the Hprt DNA sequence. al., Proc. Nat'l Acad. Sci. USA 84:5197-5209 (1987)) was follows: Plasmid PLNH2 (also known as PNHP-1, see et

To evaluate the grafted cells histologically, the rats were perfused transcardially and their brains were sectioned and stained with Nissl stain and cresyl violet for general morphological characteristics of immunocytochemical methods to establish the presence of the specific cell antiogenic markers fibroblasts and glial fibrillary acidic protein (GFAP) and GFAP (Gage et al., Exp. Neurol. 102:2-13 (1988); 1:2000 dilution) Baralle, University of Oxford, England) (1:2000 dilution) and GFRP (Gage et al., Exp. Neurol. 102:2-13 (1988); 1:1000 in TBS containing 0.25% Triton-X and 3% goat serum or with the monoclonal antibody, mouse IgG2a, against a thorough rinsing, the sections were incubated for 1 hr containing 0.25% Triton-X and 1% horse serum, followed by several rinses in TBS containing 0.25% Triton-X and 1% goat serum or 1% horse serum. The sections were then incubated for 1 hr at room temperature with a complex of avidin and biotinylated horseradish peroxidase (Vectastain, ABC kit, Vector Labs, Burlingame, CA) diluted 1:100 in 0.1 M TBS containing 0.25% Triton-X and 1% goat serum or 1% horse serum, followed by thorough rinses. The peroxidase was visualized by reacting with 0.05% NiCl₂ and 0.01% H₂O₂ in TBS for 15 min at room temperature.

Histological Analyses

Primary rat fibroblasts grafted to the neostriatum of the rat seven weeks earlier are illustrated in Fig. 6. Serial 40 mm-thick sections were stained with anti-fibronectin (A), cresyl violet (Disserry et al., Histology and London 1970), (B) and anti-GFAP (C). The surviving cells appeared to be intact and to have clumped or aggregated around the area of the infection. The cells played an intense staining for fibronectin at the core of the graft, with a clear GFAP-staining at the edges of the graft. However, little one sees with the cannula tract alone. However, little GFAP-staining was observed in the graft itself. With cresyl violet, small, round, darkly stained cells were observed in the region of the graft which could either be microglia or lymphocytes that had infiltrated the area in many of the grafts. Many of the fibroblasts could be detected response to injury. Macrophages could also be detected blasts was similar to the primary fibroblasts (not shown). Astrocyte grafts also had a similar appearance, except they were not fibronec-tin-positive, and stained for GFAP through the center of the grafts. For all three cell types, no differences were observed between the different cell types and control cells. An important feature of these cell suspension grafts is that most of the cells remained aggregated near the site of injection and did not appear, under these circumstances, to migrate very far from the injection site into the host brain. This apparent lack of migration could certainly be different for other donor cell types and graft sites, and therefore the area of the brain into which the donor cells are to be implanted, the nature of the donor cells, and

The presence of human Hprt enzyme activity demonstrated that the rat 208F cell lines grafted into the brain survived and continued to express the Hprt transgene at easily detectable levels for at least 7 weeks. Furthermore, the implanted cells could be successfully recultured, producing cells morphologically identical to the starting cultures. Infection of these cells with helper virus resulted in the production of Hprt viruses, confirming the identity of the cells and indicating that the provirus remained intact. Studies with the neo-*L*-citrulline vector confirm the survival of these cells with the citrulline selection. The expression of Hprt in the infected cells was approximately 20% of that in the uninfected cells.

Implanted cells were dissociated out and prepared for reculturing and for biochemical and molecular characterization by dissociating the cells with trypsin. For the preparation of the human Hprt activity, cell extracts were prepared from the bulk of each sample as previously described and examined by a polyacrylamide gel iso-electric focusing Hprt assay (jolly et al., Proc. Nat'l. Acad. Sci. USA) 80:477-481 (1983); Miller et al., J. Biol. Chem. 259:7842-7849 (1984); Miller et al., Science 225:630-632 (1984); Gruber et al., Science 230:1057-1061 (1985), and see et al., Gene 53:97-104 (1987)). The remainder of each sample was placed into culture. The results of an Hprt gel assay of rat 208F cells Hprt resistance after infection with the Hprt vector implanted into one side of the rat basal ganglia 3 and 7 weeks after transplantation and prior to analysis are shown in Fig. 7.

Characterization of Implanted Cell

The phenotype of the target cells for the transgene may be important factors for the selection of donor cells.

The 777 bp HgA1-PstI fragment was isolated from NGF cDNA from plasmaid PSN15, (Wolfe et al., Mol. Biol. Med. 5:43-59 (1988)). Briefly, the N1 vector described by Wolfe, supra (supplied by Dr. Harvard Medical School, 1988), under control of an internal Rous sarcoma virus promoter.

Genet. 1:327 (1982)), under control of an internal Rous insertion of transposition Tn5 (Southern et al., J. Mol. Appl. 5:43-59 (1988)). The vector also included a dominant selectable marker encoding the neomycin-resistance gene constitutively. The vector also includes a gene that is secreted to encode the precursor to NGF that is believed (Edwards et al., Nature 319:784 (1986)) and is believed to encode the shorter NGF transcript that predominates in mouse tissue receiving sympathetic innervation corresponds to the viral S⁺ LTR. This insert (1983)), under control of the viral S⁺ LTR, this insert encodes the precursor to NGF that is secreted (Nature 302: 538 (1983); Ulrich et al., Nature 303:821 HgA1-PstI fragment of mouse NGF cDNA (Scott et al., Nature 302: 538 (1983); Ulrich et al., Nature 303:821 (1982)). The PLN-BRN vector contains the 777 base pair (1982)). A retroviral vector, similar to one described previously (Varma et al., RNA Tumor Viruses; Weiss et al., Cold Spring Harbor Press, Cold Spring Harbor, NY, p. 233) was constructed from Moloney murine leukemia virus (MuLV) construct was conducted to determine whether sufficient to express foreign gene (transgene) product. The present example was conducted to determine whether sufficient to express foreign gene (transgene) product. The present when implanted into the mammalian brain and can survive genetically modified using retroviral vectors can surviveously damaged brain function.

Construction of NGF Vector PLN.BRN

The above example demonstrated that cultured cells transgene product can be made by genetic modification of the genome to complement or repair an absent or previously damaged brain function. Cells in vivo to complement or repair an absent or previously damaged brain function.

Grafting of Genetically Modified Cells Expressing NGF to the Damaged Brain

Example II

School, Harvard, Boston) was cloned into the PstI site of plasmid PSP64 (Promega, Madison, WI). Plasmid PSP65 was digested with restriction enzymes PstI and Hgal using established methods (Maniatis et al., Cold Spring Harbor, New York, (1982) and the 777 Press, Cold Spring Harbor, New York, (1982) and the 777 basepair DNA fragment containing the NGE sequences was isolated by standard purification methods (Maniatis, et al., Supra). The 777 bp fragment was then blunt ended by Klenow polymerase as described by Maniatis et al., (University of California, San Diego, CA).

in Proc. Nat'l. Acad. Sci. (USA) 84:5197-5201 (1987), ligated into plasmid PLMPL as described by Yee et al., incorporated by reference herein and supplied by Drs. Lee and Xu, (University of California, San Diego, CA).

Plasmid PLMPL was digested with Hind III to remove the metallothionein promoter and most of the Hprt DNA, and the overhangs of the 777 bp fragment isolated as above were similarly repaired. The 777 bp fragment was then blunt ended and ligated into the digested plasmid PLMPL. The result was a plasmid purified from plasmid JD204 described by de Wet et al., in Mol. Cell. Biol. 7:725-737 (1987) as follows: A 1717 bp HindIII-SspI fragment from the E. coli K12 strain was digested with PstI and a 1321 bp HindIII-SmaI fragment from plasmid JD204 and a 1321 bp HindIII-SmaI fragment derived from plasmid DSV2NEO described by Southern and Berg in Mol. Appl.

Genet. 1:327-341 (1982) were ligated with a fragment containing a mutated Rous sarcoma virus (RSV) promoter in a gene. BamHI-ClaI fragment isolated from plasmid PLMPL. BamHI-ClaI fragment isolated from plasmid JD204. Klenow base fragment was ligated to a 2.1 kilobase exon enzymes BamHI and ClaI and then digested using restriction endonucleases Klenow and SspI. The chloride centrifugation (Maniatis et al., Supra). The lashed methods for plasmid purification including cesium chloride centrifugation (Maniatis et al., Supra). The purified plasmid JD204 was digested with EcoRI and gel electrophoresis. The 777 bp fragment was then isolated as above and purified from plasmid JD204. The 777 bp fragment was then digested with Klenow and a 1321 bp fragment isolated from plasmid DSV2NEO described by Southern and Berg in Mol. Appl. Genet. 1:327-341 (1982) was ligated with a fragment containing a mutated Rous sarcoma virus (RSV) promoter in a gene. The 777 bp fragment was then digested with PstI and a 1321 bp HindIII-SmaI fragment derived from plasmid JD204 and a 1321 bp HindIII-SmaI fragment isolated from plasmid DSV2NEO described by Southern and Berg in Mol. Appl.

Plasmid PLLRN1 is depicted in Figure 8.
300 bp BamH1-Hind III fragment from plasmid PUCRH.

Plasmid PUCRH was produced as follows. Plasmid PRSneo was restricted using HindIII and the linearized plasmid was ligated with the remaining fragment of California, San Diego, CA obtained by restriction using HindIII to remove Hprt sequences. Plasmid PPR1 was obtained from Dr. Friedmann, University of California, San Diego, CA. Plasmid PPR1 (Figure 9) (supplied by Dr. Friedmann, University of California, San Diego, CA) obtained by restriction using HindIII and RsaI. The resulting plasmid was called PRH and was then restricted with PstI and PAC(-). PR(+), was then restricted with BamHI. The resulting linearized plasmid was ligated with a fragment obtained from plasmid PSVori by restriction with SalI and PstI. The resulting plasmid was termed PSVORI. Gattelersburg, MD) that had been restricted with SalI and PstI. The resulting plasmid was termed PSVORI.

The plasmid PRH+S+ that resulted from ligation of the BamHI restricted plasmid PN(+) and the BamHI restricted plasmid PSVORI was then restricted with MspI and the overhangs 5' ends were replicated using Klenow polymerase as described above. This fragment was ligated with a MspI (Bethesda Research Laboratories) linearized alkaline phosphatase (Boehringer Mannheim, W. Germany). The resulting plasmid was called pMPRH and contained the Hprt cDNA expressed from the RSV promoter.

Plasmid pMPRH was subject to site-directed mutagenesis as described by Kunkele et al., Proc. Natl. Acad. Sci. USA 82:488-492 (1985), incorporated by reference. Plasmid pMPRH was mutagenesis as described to site-directed

reference herein in order to alter the polyadenylation
signal ATTAA to AGCAA. After mutagenesis the resulting
plasmid was restricted using HindIII and the resulting
fragment was ligated to a HindIII fragment from pRSV-
SV40 plasmid PUC19 (Bethesda Research Laboratories) to
produce plasmid PUCRSV. Plasmid PUCRSV was restricted
using BamHI and PstI to produce a fragment containing the
RSV promoter. This fragment was ligated to a PstI-SstI
fragment containing the gene encoding Hprt obtained by
restriction of the plasmid PUC19, forming plasmid PUCRH (Figure 10).
Example 1, supra and to a BamHI-SstI fragment obtained
from pRSV vector pRSVneo pJD204 and the 1321 bp HindIII-SmaI
fragment from pUCRH, the 1717 bp HindIII-SspI fragment from
resulting ligation between the BamHI-HindIII frag-
ment from pUC19, after transfection and purification, the plasmid
resulting from ligation of the 6.1 BamHI-SstI base
BamHI-ClaI fragment from pUC19 plasmid PLLRNT was termed
PLN-BRN1. (Figures 11 and 12).

After transfection and purification, the plasmid
resulting from ligation of the 2.1 kilobase
fragment from pUC19 and the 2.1 kilobase
BamHI-ClaI fragment from pUC19 plasmid PLLRNT was termed
PLN-BRN1. (Figures 11 and 12).

Preparation of Transmissible Retrovirus

Transmissible retrovirus was produced by transfecting cells (Miller et al., Mol. Cell. Biol. 6:2895 (1986)), which were supplied by Dr. Miller, Fred Hutchinson Cancer Research Center, Seattle, WA), by the calcium phosphate
co-precipitation method (Graham et al., Virology 52:456 (1973)), and using medium from these cells to infect #2 (1983)) in the presence of 4 ug/ml Polybrene (Sigma Chem-ical, St. Louis, MO). Virus from the #2 clone producing units/ml, was the highest titer, a x 10⁵ colony forming units/ml, was

used to infect the established rat fibroblast cell line 208F (Quade, *Virology* 98:461 (1979) as described by Miyamoto et al., in Proc. Natl. Acad. Sci. USA (1988)). Were expanded and tested for NGF production and secretion by a two site (ELISA) enzyme immunoassay (Korsachiv et al., Proc. Natl. Acad. Sci. (USA) (80:3513-3516 (1983)), using commercial available reagents according to the manufacturer's protocol (Boehringer Mannheim, Biochemicals). Selected in medium containing the neomycin analog G418, individual neomycin-resistant 208F colonies, were expanded and tested for NGF production and secretion by a two site (ELISA) enzyme immunoassay (Korsachiv et al., Proc. Natl. Acad. Sci. USA (80:3513-3516 (1983)), as described in medium containing the neomycin analog G418, W. Germany). The culture supernatant was collected NGF contained 1.7 ng NGF/mg total cellular protein and secreted NGF into the medium at a rate of 50 pg/hr 10⁵ cells. The NGF secreted by this clone was biologically active, as determined by its ability to induce neurite outgrowth from PC12 rat pheochromocytoma cells (Green, et al., Proc. Natl. Acad. Sci. USA 73:2424 (1976)). Uninjected 208F cells, in contrast, did not produce detectable levels of NGF in either assay.

Fimbria fornix Transsection

Fimbria fornix transsection was performed as described by Gage et al., *Brain Res.* 268:27-37 (1983) and in *Neuroscience* 19(1):241-255 (1986), both of which are incorporated by reference herein. Briefly, adult female Sprague-Dawley rats (Bantin and Kingman, San Francisco, CA) weighing between 200g and 225g at the beginning of the experiment were used. The animals were anesthetized with intraperitoneal injections of a ketamine-xylazine mixture (10 ug/kg Rompun, Hoechst, Park-Davis Ann Arbor, MI, and 5 ug/kg Rompun, Hoechst, Frankfurt, W. Germany). Unilateral aspiration (10 ug/kg Ketalet, Park-Davis Ann Arbor, MI, and 5 ug/kg Rompun, Hoechst, Frankfurt, W. Germany). Unilateral aspiration (10 ug/kg Ketalet, Park-Davis Ann Arbor, MI, and 5 ug/kg Rompun, Hoechst, Frankfurt, W. Germany).

Assay for NGF Production and Secretion

Miyamoto et al., in Proc. Natl. Acad. Sci. USA (1988)). 208F (Quade, *Virology* 98:461 (1979) as described by Miyamoto et al., in Proc. Natl. Acad. Sci. USA (1988)).

At 2 weeks following surgery the rats were perfused and their brains were removed, fixed overnight and placed

Immunohistochemistry

of suspended cells were injected free-hand using a pipette to the cavity in the animals. A piece of gel foam was gently placed on the surface of the cavity and flattened into the cavity in the animals. A piece of cellulose sponge was inserted into the cavity and later ventricile hamilton syringe into the cavity and later ventricile and resuspended in complete PBS at 10⁵ cells/ml. Four ml and 5% for 4 min. At 4°C, washed twice with complete PBS, x 5 for 4 min. Cells were pelleted by centrifugation at 1000 rpm. Cells were serum to inactivate the CAC12 (complete PBS) and 5% rat serum to inactivate the complement with 1 mg/ml glucose, 0.1 mg/ml each MGCl2 and and 1 mM EDTA and taken up by trituration with PBS and phosphate buffered saline (PBS) containing 0.05% trypsin cells were removed from confluent plates with Dulbecco's retrovirus-infected (NGF secreting) and control 208F

grafts (Fig. 13C-F). Infected cells than in animals that had received control medial septum in animals that had received grafts of number of remaining neurons on the lesioned side of the vival of cholinergic cells bodies indicated a greater for choline acetyltransferase (CHAT) to evaluate the similarity in both groups (Fig. 13A,B). Sections stained comparable in both groups (Fig. 13A,B). Sections stained cific marker, revealed robust graft survival that was sections stained for fibronectin, a fibroblast spe-

ring 8 received uninjected control cells. rats received grafts of infected cells while the remainin fornix lesions as described above were made in 16 rats; 8 the same complete unilateral aspiration lesion. Fimbra All animals in each of the experiments received as described in Gage et al., Brain Res. 268:27-37 (1983). to inflict a partial denervation on the hippocampus target, hippocampus as well as the overlying cingulate cortex to

tions 40 μm thick were cut on a freezing slide
 in phosphate-buffered 30% sucrose for 24 hr at 4°C. Sec-
 tion was labelled immunohistochemically by standard
 procedures using polyclonal antibodies to fibronectin to
 evaluate fibroblast survival. Polyclonal antibodies to
 cholinergic acetyltransferase (anti-CHAT antisera) were also
 generated to evaluate the survival of cholinergic cell
 bodies as described by Gage et al. in J. of Comparative
 Neurol. 269:147-155 (1988), incorporated by reference
 herein. Tissue sections were processed for immunohistochemistry
 according to a modification of the avidin-
 biotin labelling procedure of Hsu et al., 29:1349-1353
 (1981), incorporated by reference. This procedure con-
 sists of the following steps: 1) overnight incubation
 for 1 hr with biotinylated goat antirabbit IgG (Vector
 body was diluted 1:1,500 with 0.1 M Tris-saline contain-
 ing 1% goat serum and 0.25% Triton X-100; 2) incubation
 for 1 hr with biotinylated goat antibody (i.e.,
 preimmune serum or absorbed antibody). The anti-
 cholinergic complex (Vector Laboratories, CA) diluted 1:200 with
 Tris-saline containing 1% goat serum: 3) 1 hr incubation
 with ABC complex (Vector Laboratories) diluted 1:100 with
 Tris-saline containing 1% goat serum: 4) treatment for
 15 min with 0.05% 3,3'-diaminobenzidine (DAB), 0.01%
 hydrogen peroxide and covered with permount and
 buffer. Immunolabelled tissue sections were mounted onto
 glass coverslips. Two sections stained for CHAT through
 the septum, 200 μm apart were used to evaluate the extent
 of cholinergic cell survival. All the CHAT-positive
 cells in the ipsilateral septum and in the contralateral
 area were counted separately and sized for planar area
 using an Olympus Que-2 image analysis system. Tissues
 were also stained for acetylcholinesterase (AChE) as
 were also stained for acetylcholinesterase (AChE) as

described by Hedreen et al., J. Histochem. Cytochem. 33:134-140 (1985), incorporated by reference herein, to evaluate the completeness of the fibrilla formix transection. Neuronal survival was quantitated (Fig. 14) and, when expressed as a percentage of the remaining cholinergic cells in the septum ipsilateral to the lesion relative to the intact contralateral septum, was shown to be 92% in animals grafted with NGF-secreting cells but only 49% in animals grafted with control cells. Previous results from the control group are comparable to previous observations in lesioned animals that had received no grafts (Gage et al., Neurosci. Res. 8:2155 (1986); Neuroscience 19:241 (1986); Heftei, J. Neurosci. USA 83:9231 (1986); Kromer, Science 235:214 Acad. Sci. 235 (1987); Gage et al., Comp. Neurol. 369:147 (1988)).

In addition to the significant increase in the percentage of ChAT-positive cells in the NGF group, these findings also showed an increase in acetylcholinesterase staining in the dorsal lateral septum, response in the dorsal lateral septum, with the most intense staining abutting the cavity containing the above results demonstrate the feasibility of continued transgene expression by cells grafted to the CNS and also present the first demonstration of a phenotype, grafted, genetically modified cells.

The above results demonstrate the feasibility of continued transgene expression by cells grafted to the CNS and also present the first demonstration of a phenotype, grafted, genetically modified cells.

structured by inserting DNA encoding the retinoblastoma gene of plasmid pGEN1-4.5Rb old (pGEN1-4.5Rb old was constructed with the 3.5 kb fragment obtained after restriction digesting the Hprt gene. The remaining plasmid DNA was enzymes HindIII and HpaI, and removing the fragment containing the Hprt gene. (obtained as described above in Example II) with the plasmid pLRB1 was obtained by digesting plasmid pLMTP1 and pLTHNL were ligated together to form pLTHNL.

fragments from three plasmids: pLRB1, pTH54 and gene transcribed from an internal RSV promoter (Figure 16). From the 5' LTR sequence and contained a neomycin-resistor (Mo-MLV) derived retroviral vector, was constructed expressing the rat cDNA for tyrosine hydroxylase (TH) (Mo-MLV) derived retroviral vector, a Moloney Leukemia virus The vector pLTHNL, a Moloney Leukemia virus

Construction of Retroviral Vector pLTHNL

The strategy for enabling fibroblasts to produce L-DOPA used in this example is based upon the ability of the enzyme tyrosine hydroxylase (TH) to catalyze the conversion of tyrosine to L-DOPA; the rate-limiting step in catecholamine synthesis. Tetrahydrobiopterine (H4-B), the co-factor for TH is required for the enzymatic activity. Since the brain contains significant levels of biopterin, and fibroblasts can reduce biopterin to H4-biopterin, TH should be active in fibroblasts situated within the brain.

This example was undertaken to demonstrate that the methods of the present invention for genetic modification of donor cells and grafting of the cells into the CNS can significantly ameliorate the signs of disease in an animal model, such as a rat model of Parkinson's disease.

Grafting of Genetically Modified Fibroblasts Expressing L-DOPA Into The CNS Of A Rat

Example III

gene (RB) intoplasmid PGEM1, available from Promega, Madison, WI, and was supplied by Dr. Lee, University of California, San Diego, CA) using HindIII and ScaI. The resulting plasmid was named pLRBL.

A 1688 bp fragment containing rat TH cDNA was obtained from the plasmid pTH54 (O'Malley, J. Neurosci. Res. 60:3-10 (1986), supplied by Dr. O'Malley, Washington University, St. Louis, MO) by digestion with BamHI and SphI.

The vector pL2RNL containing the retroviral provirus for plasmid pL2RNL (described above in Example I) to obtain the vector pLTHNL containing the retroviral provirus for pLTHNL is shown in Figure 17.

The derivation of and circular restriction map for pLTHNL is shown in Figure 17.

The gene encoding the enzyme tyrosine hydroxylase, ing the gene encoding the enzyme tyrosine hydroxylase, was cloned into producer cells to produce virus carrying transfection into 223-232 (1977), incorporated by reference herein, into the amphotropic P317 helper line supplied by Dr. Miller, Fred Hutchinson Cancer Research Center, (Dr. Miller, Fred Hutchinson Cancer Research Center, Seattle, WA). Two days post-transfection, media from these cells were filtered and used to infect the G-418-resistant #2 clone (#2/TH) that contained the high-level of TH activity and produced the highest titer of virus (5×10^5 /ml) was selected. Immortalized, rat fibroblasts (208F) (Quade, Virology 98:461-465 (1979)) were infected at a multiplicity of infection (MOI) of less than 10⁴ with LTHNL virus produced by the #2/TH.

Cells were infected with a C1a1 and small fragment obtained from plasmid pL2RBL (described above in Example I) to obtain helper-free retrovirus was produced and retroviral infections were done as described by Wigler et al., Cell 11:223-232 (1977), incorporated by reference herein, into 2895-2902 (1986), supplied by Dr. Miller. A ecotropic #2 helper line (Miller et al., Mol. Cell. Biol. 6:2895-2902 (1986), supplied by Dr. Miller). A

cultured cells were scraped off plates as in the assay of tyrosine hydroxylase activity except ascorbic acid (final concentration of 50 μ M) was added to the cell pellets prior to freezing. Cells were grown in DME plus 10% fetal calf serum. Cells were cultures were supplemented with 0.1 mM GMPA 16 hours prior to harvesting. Cells were homogenized in 250 μ l of ice cold 0.1 N HClO₄/0.1% sodium metabisulfite/0.2% Na₂EDTA containing 5 ng/ml of 3,4-dihydroxybenzylamine (DHB) as internal standard. Dopamine DOPA, 3,4-dihydroxyphenylacetic acid (DOpac),

Assays of Catecholamines and their Metabolites

Contaminating calcium or magnesium chloride and the cells were scraped off the plates. The cells were homogenized in 0.15 ml of ice cold 50 mM Tris/50 mM sodium pyro-phosphate/0.2% Triton X-100, adjusted to pH 8.4 with acetic acid, and were centrifuged at 32,000 x g for 15 min at 2-4° C. The supernatant fraction was used for both the and protein measurements. The activity was measured with (1986), but with ^{14}C -labelled 20 mM tyrosine, 1 mM 6-methyl-5,6,7,8-tetrahydropterin (6MHP \ddagger) (Calbiochem, La Jolla, CA), and potassium phosphite buffer (pH 6). Protein was determined by the method of Lowry et al. (J. Biol. Chem., 193:265 (1951)) using bovine serum albumin as standard.

Assay of Tyrosine Hydroxylase Activity

producer cells. G-418 resistant clones were established for further study. All retroviral infections were done in the presence of 4 µg of polybrene (Sigma) per ml. Cells were selected for expression of the neomycin-resis-

and DHA were extracted from the supernatant fraction by alumina absorption (Anton et al., J. Pharmacol. Exp. Ther. 138:360-375 (1962)) and eluted with 150 μ l of 0.1N H₃PO₄. They were analyzed by HPLC with electrochemical detection as described by Iovone et al., Brain Res. 418:314-324 (1987), with the mobile phase modified to contain a higher concentration of sodium octylsulfate (0.45 M) and lower pH (2.8). Homovanillic acid (HVA) was analyzed by HPLC. The concentration of catecholamines and metabolites in the media was also determined using a different method, HPLC-EC. The alumina extraction procedure described above was omitted and the media was adjusted to 0.1 perchloric acid/0.01M EDTA was centrifuged 10,000g x 10 min to remove precipitated material and used directly for HPLC-EC. In this system, the whole phase consisted of 0.137% SDS in 0.1 M phosphate buffer, pH 3.2 (Buffer A) or 40% methanol in 0.1 M phosphate buffer, pH 3.05 (Buffer B). Compounds in sample were eluted for 12 minutes in 100% Buffer A, followed by a gradient increasing linearly over 30 minutes to 100% Buffer B. The eluent was then passed through a series of 16 coulometricic electrodes set at 60 mV increments.

Female Sprague-Dawley rats received a unilateral injection of 12 μ g in 2 μ l saline-ascorbate 6-hydroxy-dopamine (6-OHDA) into the medial forebrain bundle (coordinates: AP=-4.4; ML=1.1; DV=7.5). Completeness of the lesion produced was assessed 10 to 20 days postinjection by either apomorphine (0.1 mg/kg, s.c.) induced rotational behavior (ungerstedt and Arbuthnott, Brain Res. 24:485-493 (1970)). Prior to transplantation, each animal was tested at least twice on separate days to establish the baseline rotatory response to apomorphine or

Rat Model of Parkinson's

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Grafted rats were tested for rotational asymmetry 1
and 2 weeks following fibroblast grafting.

Post-Grafting Behavioral Testing

Received injections of noninjected fibroblasts.
2 mm area at each site. Control lesions animals
of 4 μ l were delivered in two equal deposits over a 1 to
 $ML=3.0$; $DV=3.5/4.5$) of the derivate caudate. A total
 $AP=2.5$; $ML=1.5$; $DV=3.5/4.5$) and caudal areas ($AP=0.4$;
rostro (coordinates: $AP=1.4$; $ML=2.0$; $DV=3.5-5.5$ to
stereotaxically into 2 to 3 separate locations within the
suppl. 522:29-37 (1983)), suspended cells were injected
all., Brain Res. 297:53-61 (1984); Dunnert et al., Scan-
cial for recovery from rotational asymmetry (Herrera et
per μ l. Since graft placement has been shown to be cru-
resuspended in complete PBS at a density of 80,000 cells
plete PBS using centrifugation at 1000 x g and were
vate the trypsin. The cells were washed twice with com-
mg/ml $CaCl_2$ (complete PBS) plus 5% rat serum to inacti-
plemented with 1 mg/ml glucose, 0.1 mg/ml $MgCl_2$ and 0.1
PBS containing 0.05% trypsin and pipetted up in PBS sup-
noninjected fibroblasts were loosened from the plates in
confuent 10 cm plates of cultured TH-injected or

Grafting of fibroblasts

males.
tion was compared in the 4 experimental groups of ani-
mals.
umber of rotations from baseline to post-transplanta-
tions/min towards the side of the lesion following
amphetamine tested). The average percent change in the
amphetamine administered; 19 apomorphine tested, 14
rotations/min towards the side of the lesion following
amphetamine administration and at least 7 ipsilateral
study (at least 7 contralateral rotations/min following
J. Neurochem. 38:737-748 (1982)) were included in the
of more than 7 turns per minute (Schmidt et al.,
amphetamine for each animal. Animals turning at a rate

with LTHRN1 virus produced by the #2/TB producer cells
immortalized, rat fibroblasts (208F) were infected

Levels of TB

Establishment of a Fibroblast Clone Expressing High

positive grafts.

tions were assessed for size and placement of fibroblast H202 in TBS for 15 min at room temperature. Mounted sec-
Chemical Co., St. Louis, MO) and 0.05% Nicl₂ and 0.01%
3,3-diaminobenzidine tetrahydrochloride (DAB) (Sigma
peroxidase was visualized by reacting with 0.05%,
1/6 goat serum, followed by thorough rinses. The
diluted 1:100 in 0.1 M TBS containing 0.25% Triton-X and
(Vectastain, ABC kit, Vectored Labs, Burlingame, CA)

plex of avidin and biotinylated horseradish peroxidase
then incubated for 1 hr at room temperature with a com-
ing 0.25% Triton-X and 1/6 goat serum. The sections were
1/6 goat serum, followed by several rinses in TBS contain-
diluted 1:200 in 0.1 M TBS containing 0.25% Triton-X and
with biotinylated goat anti-rabbit IgG (Vectastain)

TBS containing 0.25% Triton-X and 3% goat serum. After
thorough rinsing, the sections were incubated for 1 hr
1:600 or polyclonal anti-fibronectin diluted 1:2000 in
polyclonal antibodies to tyrosine hydroxylase diluted
sections were incubated for 24 hrs at 4°C with rabbit
(TBS) solution (pH 7.4) containing 0.25% Triton-X. The
briefly, the sections were rinsed in Tris-buffered saline
tyrosine hydroxylase antibody (EugeneTech, New Jersey).

violet, fibronectin (FB) or TH using a polyclonal anti-
microtome. Alterнате sections were stained for cresyl
for 48 hrs and then sectioned (40 μm) on a freezing
brains were postfixed overnight, placed in 30% sucrose
deeply anesthetized and perfused with 10% formalin.
Following the final behavioral test, rats were

Histological Methods

CELS

Cell extracts from the 208/T-H-8 fibroblasts express-
ing TH and control 208F fibroblasts were assayed for
L-DOPA (Table 2). Only 287F/T-H-8 cells cultured in media
supplemented with 6MPA produced L-DOPA. Control cells
did not contain any detectable amounts of L-DOPA.
Dopamine and its metabolites DOPA and HVA were below
detectable levels in both 208F/T-H-8 and 208F/control

Fibroblasts Expressing TH Produce and Secrete L-DOPA.

*TH activity is expressed in units of pmoles DOPA/mg protein

TH Activity*	Cel1 Line	TH Activity*	Cell Type
208F/TH-8	1.7	208F/TH-11	2.6
208F/TH-9	2.9	208F/TH-12	0.4
208F/TH-11	0.0	208F/CONTROL	0.8
			Rat Striatum

and 12 G-418-resistant clones were established. Table I shows the TH activity of 3 of these 12 G-418 resistant clones with the highest TH activity and the TH activity of the 42 producer line. The TH activity of the clones with the highest activity (clones 208F and TH-8) contained approximately a quarter of the TH activity of rat striatum. The 208F/TH-8 clone that contained the highest TH activity, was chosen for further study.

excluding those from statistical analyses. The immunoreactivity data from the rats with nonsurviving grafts were staining to the syringing tract (Figure 18C and D). Behavior non-surviving based on the confinement of fibronectin and B). Only 4 out of 31 grafts were classified as large in size regressed placement (Figure 18A due to large positive grafts were typically moderate fibronectin positive grafts were regenerated capsule. Survival to many areas within denervated caudate. Survival to grafts survived intraparenchymal transplants fibroblast grafts.

Histologic examination of grafts

MHPG or HVA in the media. TH-infected media. There was no detectable DOPA, DA, control 208F media and 239 ng/hr per 10^6 cells in media of the 208F/TH-8 cells: 63 ng/hr per 10^6 cells in media shown in Table 2, L-DOPA was also detected in the TH-infected media. There was no detectable DOPA, DA, MHPG or HVA in the media.

1. L-DOPA concentration is expressed in units of nanograms (ng)/mg protein for cell extract and in units of ng/hr/ 10^6 cells for cell media.
2. Cells incubated in normal media.
3. Cells incubated overnight in normal media supplemented with 0.1 mM DL-6-Methyl-5,6,7,8-tetrahydroxypterin.
4. N.D. not determined

Cell Clone		Cell Extract		Cell Media	
L-DOPA Concentration of Cell Extracts and Media	L-DOPA Concentration of Cell Extracts and Media	no 6MPH ₄ ² + 6MPH ₄ ³	no 6MPH ₄ ² + 6MPH ₄ ³	<0.25	<0.25
208F/CONTROL	no 6MPH ₄ ² + 6MPH ₄ ³	<0.25	<0.25	N.D.	63
208F/TH-8	no 6MPH ₄ ² + 6MPH ₄ ³	<0.25	<0.25	1.38	239

Table 2

These results demonstrate that the rat cDNA coding for the TH gene can express functional TH enzymatic activity when transduced into the rostral caudate region, they subsequently produce detectable levels of L-DOPA in fibroblasts do not produce detectable levels of L-DOPA in vitro and do not attenuate the rotational asymmetry of these rats, the ability of these DOPA-producing cells to attenuate these rat's rotational symmetry must be due solely to the presence of the TH gene within the cells.

These data demonstrate an effect on rotational behavior for at least two weeks, since we wanted to correlate these data with histological analysis.

The number of drug-induced rotations 2 weeks following transplantation (Fig. 19 bottom) was dependent on graft placement. Rats with fibroblast grafts confined to cauda striatum (Fig. 19 top) (AP=0 to 0.4) had no significant changes in rotational behavior. Rats which had significant changes in rotational behavior. Rats which had survived TH-injected fibroblasts in rostral caudate striatum (AP=1.4 to 2.2) showed an average 33% reduction in drug-induced rotations 2 weeks following transplantation (Fig. 19 bottom). When these DOPA-producing fibroblasts were expressed in the TH gene can produce and secrete L-DOPA in vitro and significantly reduce the rotational asymmetry implanted into the rostral caudate region, they subsequently implant into the rat model of Parkinson's. Since control rats do not produce detectable levels of L-DOPA in vitro and do not attenuate the rotational asymmetry of these rats, the ability of these DOPA-producing fibroblasts were

The number of drug-induced rotations for each individual animal were compared before and 2 weeks after transplantation. Rotations from rats tested with amorphine were pooled with those from rats tested with amphetamine since no difference was seen between these groups.

Effect of Grafts on Rotational Asymmetry

was not observed in the fibroblasts either in vivo or in vivo.

The effect of the DOPA-producing fibroblasts on rotatiorial behavior were dependent on placement in the rostral caudate. Previous data utilizing fetal neural grafts into rats have shown that attenuation of rotation is best achieved when the grafts are placed into the rostral caudate, Dunnett. Since the tional asymmetry is best achieved when the grafts are placed into the rostral caudate, Dunnett. Since the fibroblasts used cannot sprout axons, the location of the graft is even more criticaly dependent upon proper graft placement.

The exact mechanism by which the DOPA-producing fibroblasts reduce rotational asymmetry remains to be determined. Presumably, once L-DOPA is secreted, there remains enough catalyst DOPA decarboxylase activity, even

The exact mechanism by which the DOPA-producing fibroblasts reduce asymmetry remains to be determined. Presumably, once L-DOPA is secreted, there within enough cathepsin D to degradate DOPA decarboxylase activity, even within these totally denervated animals (Lloyd et al., 1970; Horngikiewicz, British Med. J. 29:172-178 (1973)), to convert L-DOPA to dopamine that then modifies drug-induced rotational behavior.

This postulated mechanism of action of these DOPA-producing cells would be consistent with the well established efficacy of systemic L-DOPA therapy for Parkinson's disease (Calne, N. Eng. J. Med. 310:523-524 (1984)). These DOPA-producing fibroblasts are in effect small localized pumps of L-DOPA.

The ability, demonstrated in this example, to modify cells to produce L-DOPA broadens the search for the ideal type of cell for transplantation therapy of Parkinson's. Any cell that can be genetically-modified to express the gene and that can survive long-term in the brain with-out forming a tumor or causing other damage, may be used. Although these particular immortalized rat fibroblasts have not formed tumors for up to three months, primary cells such as primary fibroblasts or primary glial cells may offer the theoretical advantage of decreased propensity for tumor formation. In addition, the use of the site for tumor formation, the patients own primary cells for an autologous graft would

It is apparent that many modifications and variations of this invention as set forth above may be made without departing from the spirit and scope of the present invention as defined by the terms of the appended claims.

These results demonstrate that a fibroblast can be genetically-modified to supply a function normally supplied by a neuron, therefore not requiring the use of fetal tissue for neural transplantation. The ability to combine transplantation modalities with gene-transfer presents a powerful method for the treatment of CNS dysfunctions. The methods of the present invention may thus be used for treatment of other models of animal and human brain disease.

For example, does offer the advantage of having large amounts of immortalized cells readily available. The chance of graft rejection, however, the use of immortalized cells used in this example, decreases the chance of graft rejection. However, the use

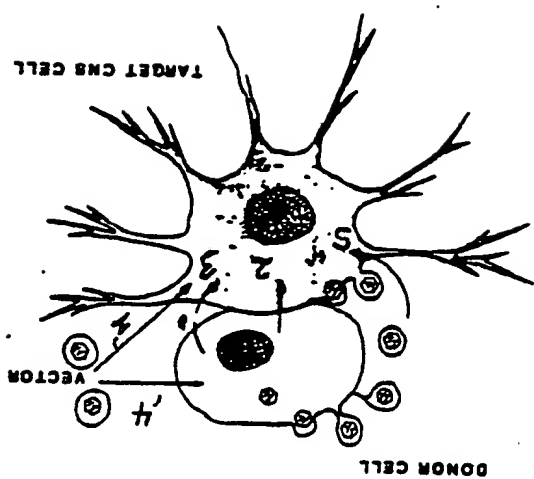
1. A method for treating defects, disease or damage to cells in the central nervous system comprising grafting donor cells into the central nervous system, said donor cells genetically modified so as to produce a molecule that directly or indirectly exerts an ameliorative effect on said cells.
2. The method of claim 1 wherein the step of grafting said donor cells comprises introducing said donor cells into the brain of a subject.
3. The method of claim 1 wherein the step of grafting said donor cells comprising introducing said donor cells into the spinal cord of a subject.
4. The method of claim 2 or 3 wherein said introducing and intravenous injection of a therapeutic transgene into said cells are modified by insertion of a therapeutic gene into the donor cells.
5. The method of claim 1 wherein said donor cells are modified by insertion of a therapeutic gene into the donor cells.
6. The method of claim 5 wherein said step of insertion comprises inserting a vector carrying said transgene, said vector being a retroviral, and neorotropic virus.
7. The method of claim 6 wherein said vector is a herpes virus vector.
8. The method of claim 6 wherein said vector is a rabies virus vector.
9. The method of claim 6 wherein said vector is a retroviral vector.

We claim:

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10. The method of claim 9 wherein said retroviral vector is the retroviral vector PLN.BRN1 having a final construction as shown in Figure 1.
11. The method of claim 9 wherein said retroviral vector is the retroviral vector PLN.BRN1 having a final construction as shown in Figure 6.
12. The method of claim 5 wherein said nonviral physical transfection of DNA encoding a transgene.
13. The method of claim 12 wherein said nonviral physical transfection of DNA encoding a transgene.
14. The method of claim 5 wherein said step of insertion into donor cells comprises electroporation.
15. The method of claim 5 wherein said step of insertion into donor cells comprises chemical mediated transfection.
16. The method of claim 15 wherein said chemically mediated transfection comprises calcium phosphate.
17. The method of claim 5 wherein said step of insertion into donor cells comprises liposomal mediation.
18. The method of claim 1 wherein said molecule is selected from the group consisting of growth factors, enzymes, gangliosides, antibiotics, neurotransmitters, neurotransmitters, neurohormones, toxins, neurite promoting molecules, anti-inflammatories, nerve growth factors, and nerve growth factor.
19. The method of claim 18 wherein said molecule is metabolites and precursors of said molecules.

21. The method of claim 18 wherein said molecule is L-DOPA.
20. The method of claim 18 wherein said molecule is tyrosine hydroxylase.
22. The method of claim 1 further comprising co-administration of a therapeutic agent for treating said disease or damage to the central nervous system.
23. The method of claim 22 wherein said therapeutic agent is selected from the group consisting of growth factors, gangliosides, antibiotics, neurotrophins, antimitabolites, neurotransmitters, precursors of these agents, and molecules and precursors of these agents.
24. The method of claim 22 wherein said therapeutic agent is cellular matter.
25. The method of claim 24 wherein said cellular matter is selected from the group consisting of adrenal chromaffin cells, fetal brain tissue cells and placental chromaffin cells, fetal adrenal matter, and material to facilitate reconnection or ameliorative interactions to the site of said damage or disease, material of injured neurons.
26. The method of claim 1 further comprising implanting material to the site of said damage or disease, material to facilitate reconnection or ameliorative interactions to the site of injured neurons.
27. The method of claim 26 wherein said material is selected from the group consisting of homogenate of brain, homogenate of placenta, whole cells, synthetic material, neurite promoting extracellular matrix, and geneticaly modified donor cells.
28. The method of claim 1 wherein said donor cells are selected from the group consisting of fibroblasts, neutrons, glial cells, keratinocytes, hepatocytes, ependymal cells and chromaffin cells.

FIGURE 1



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FIGURE 2

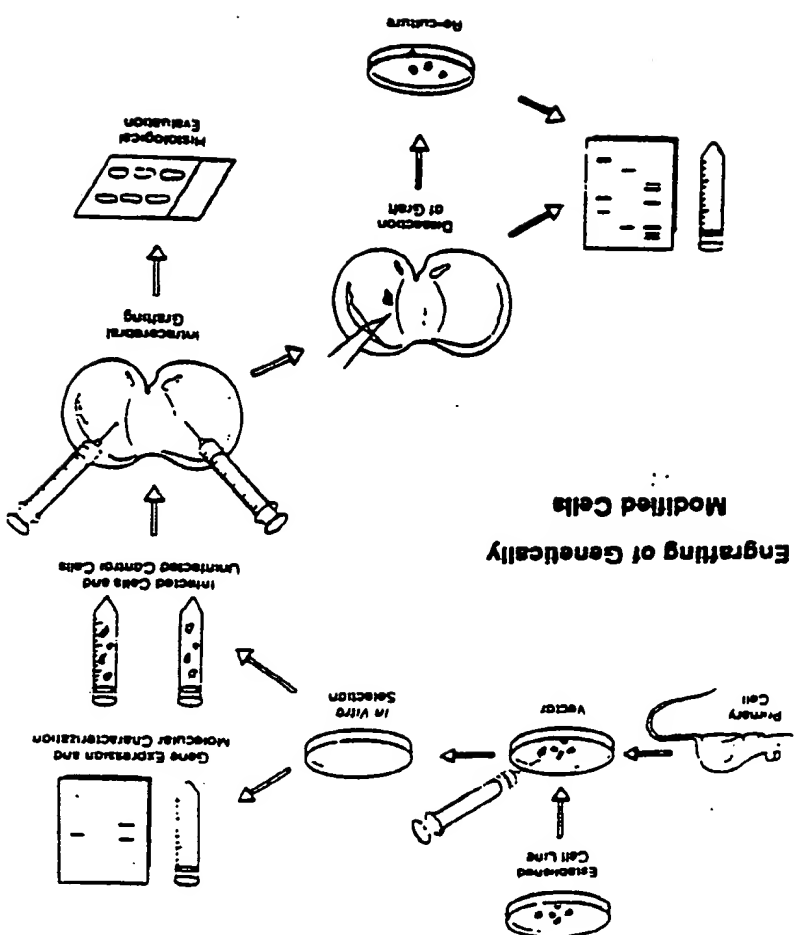


FIGURE 3

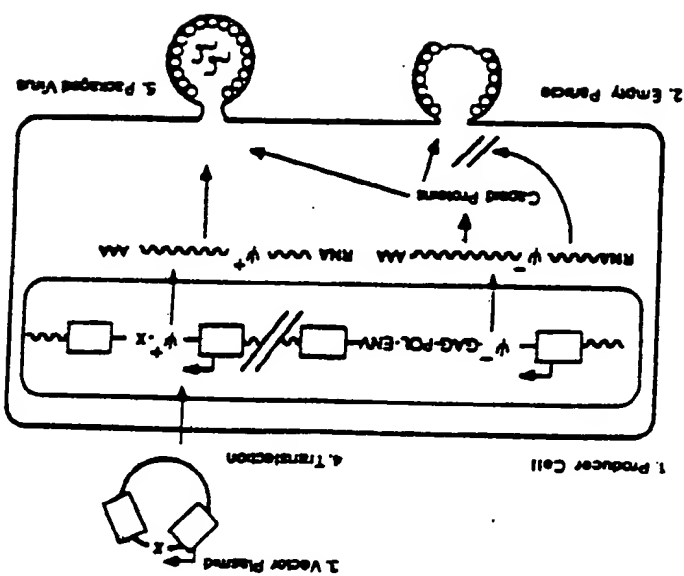
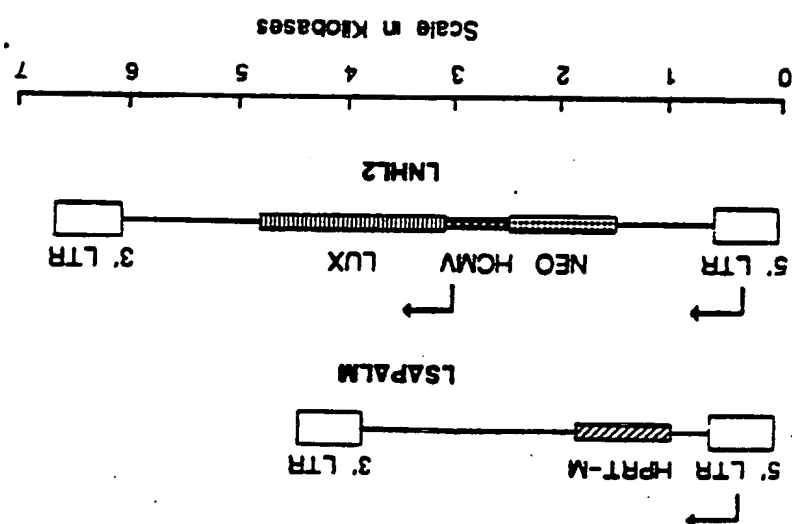


FIGURE 4



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FIGURE 5

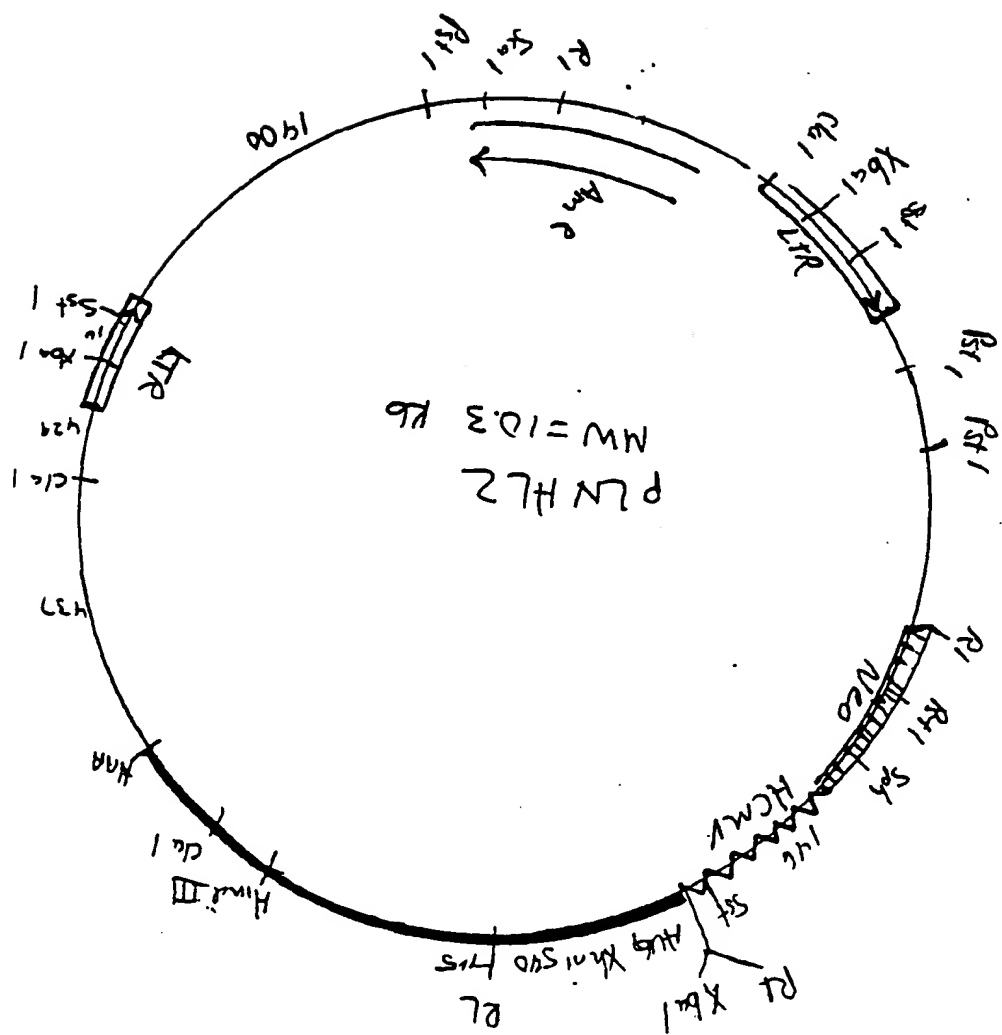
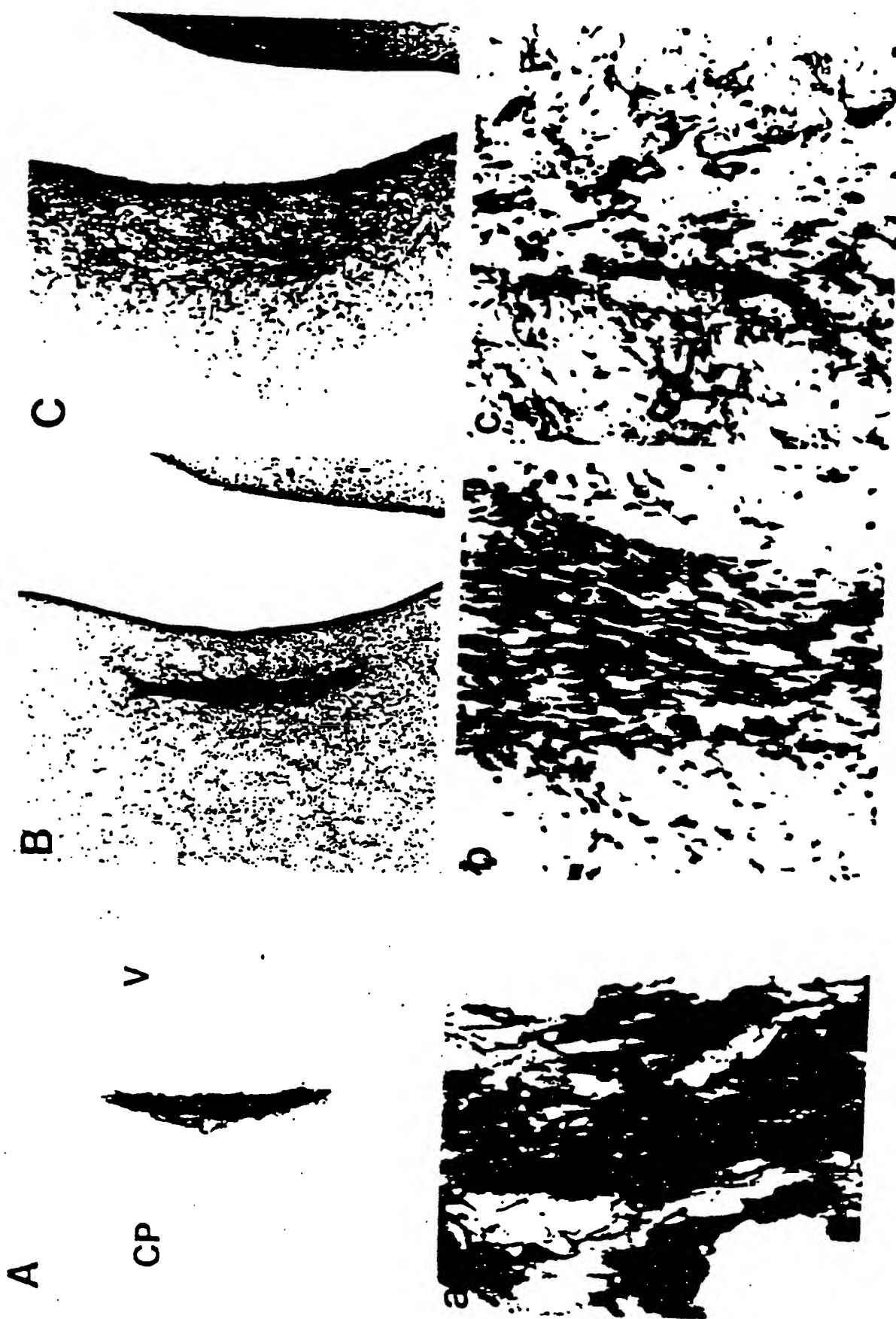


FIGURE 6



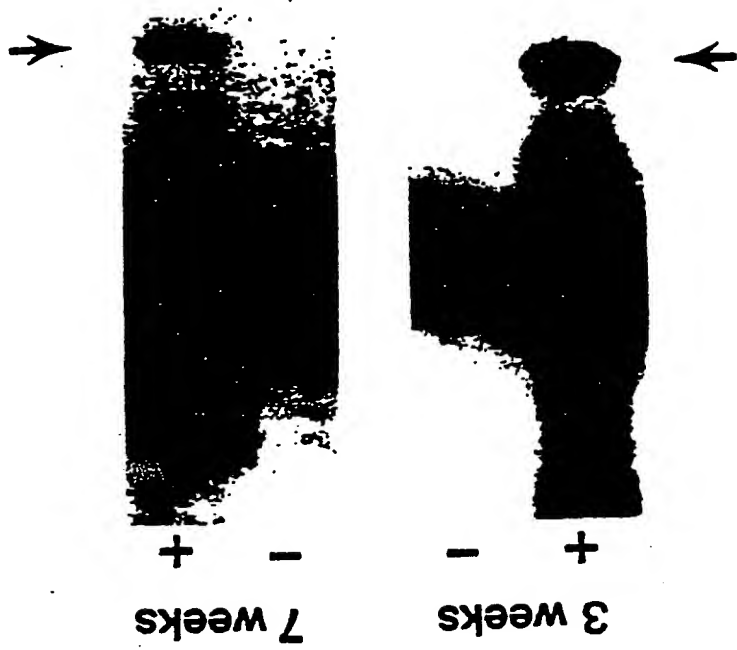
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FIGURE 7

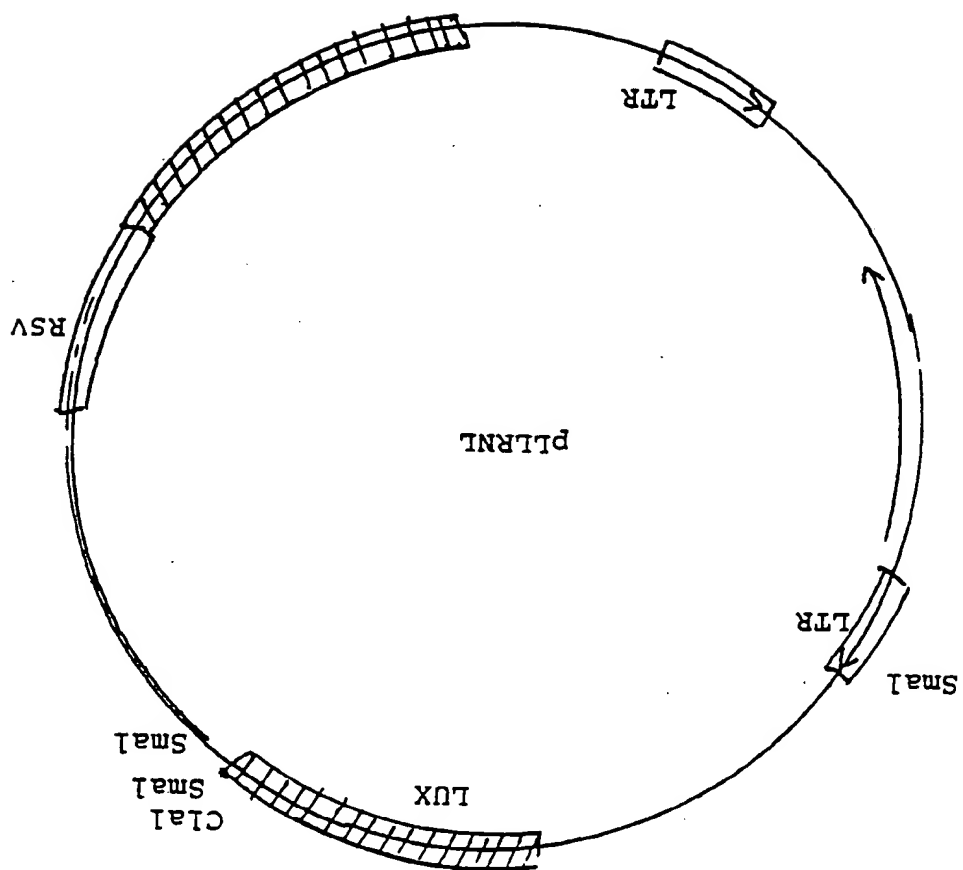


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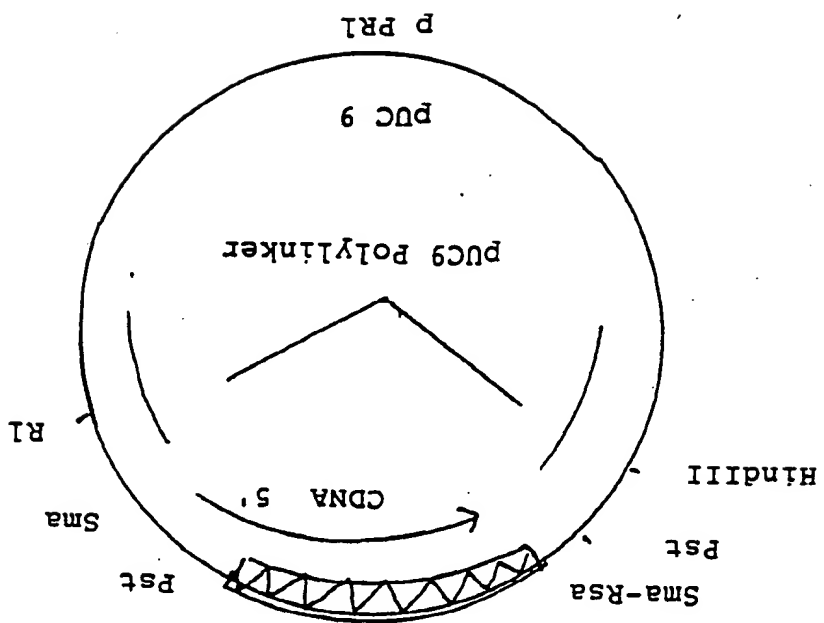
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FIGURE 8



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FIGURE 9

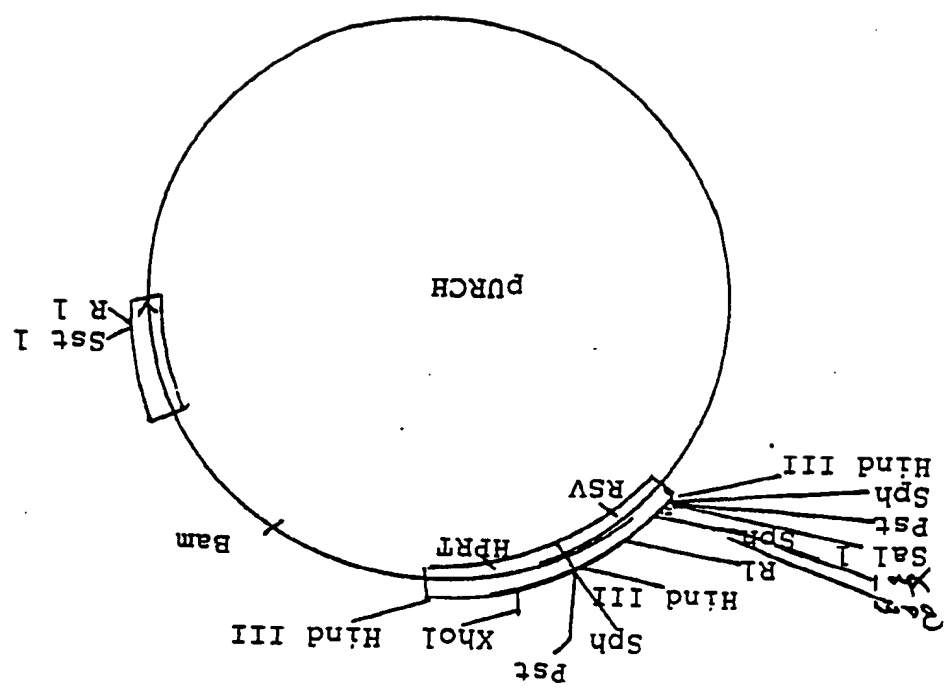


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FIG 10

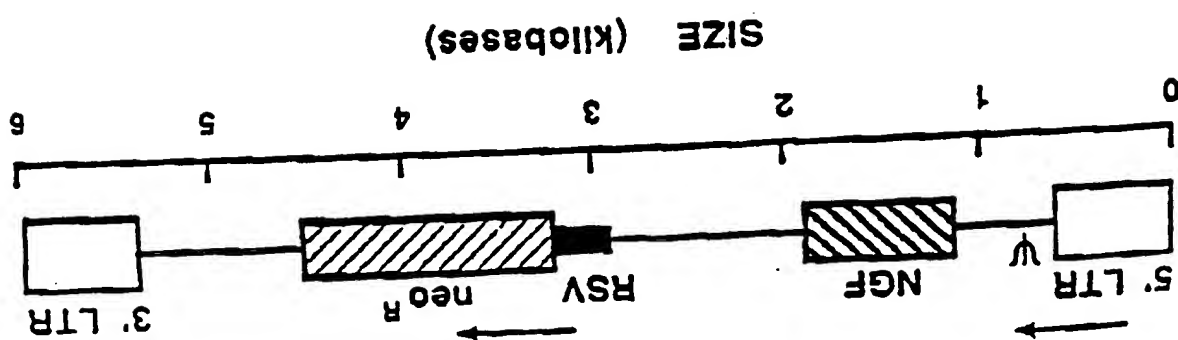


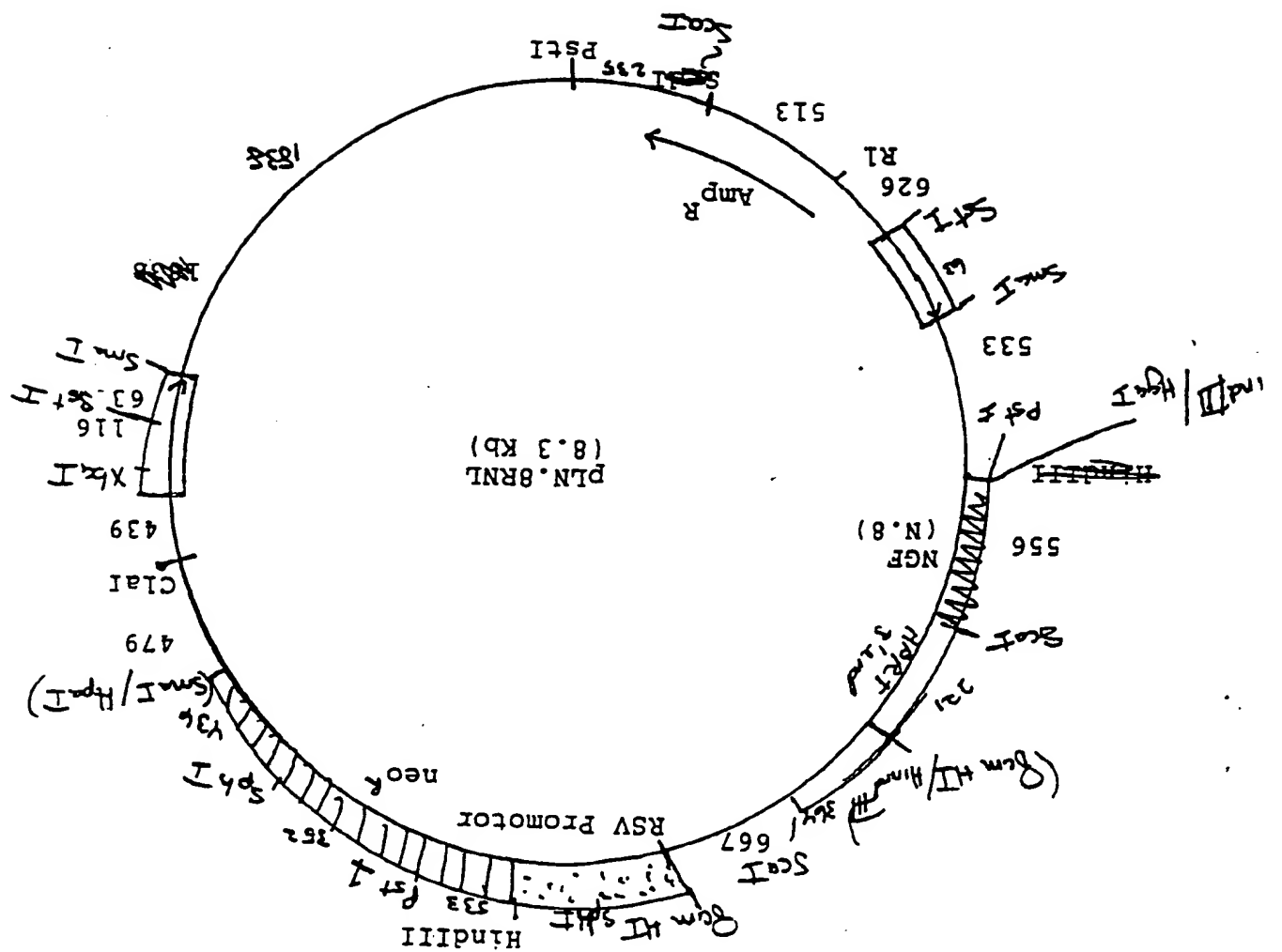
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FIGURE 11





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FIGURE 13



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FIGURE 14

Number of ChAT-IR Cells

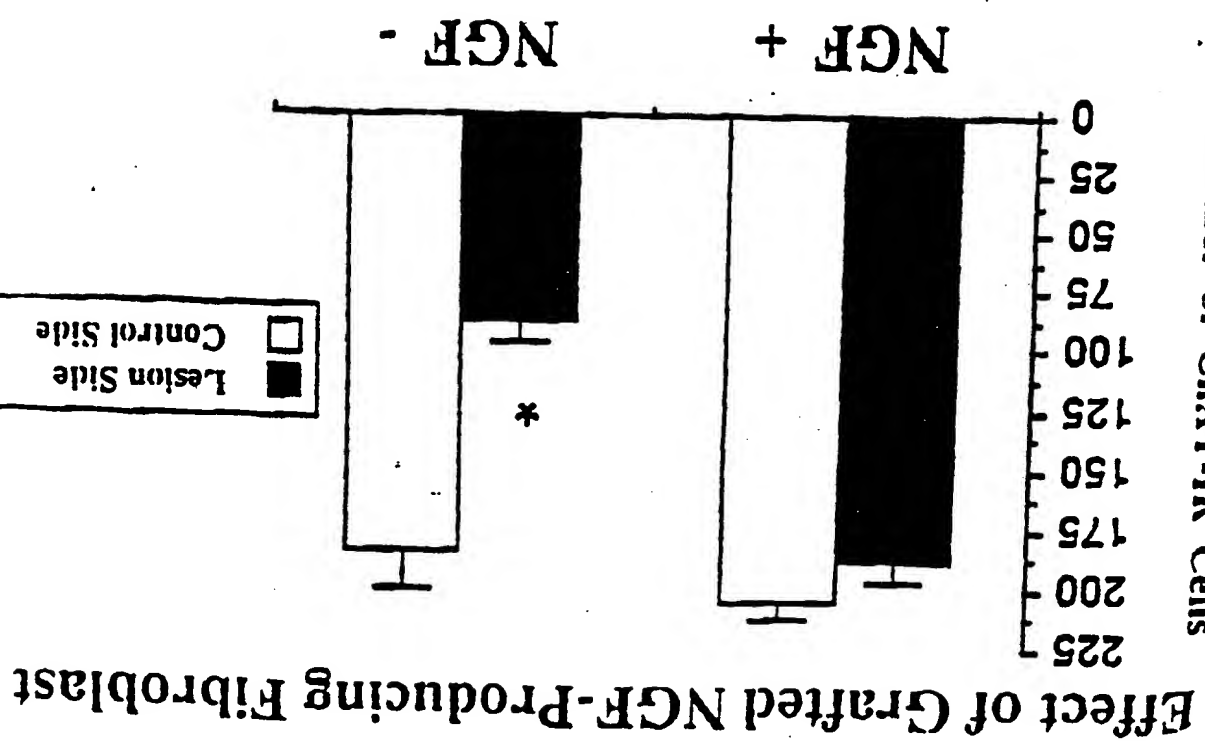
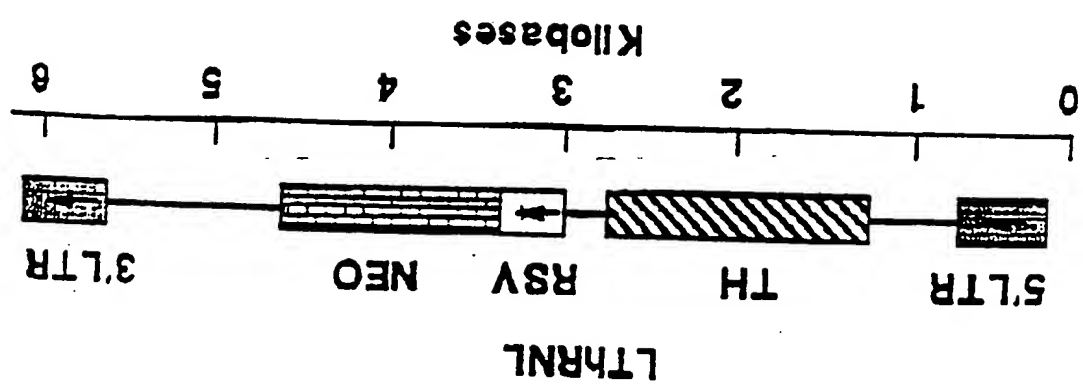


FIGURE 15



FIGURE 16



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FIGURE 17

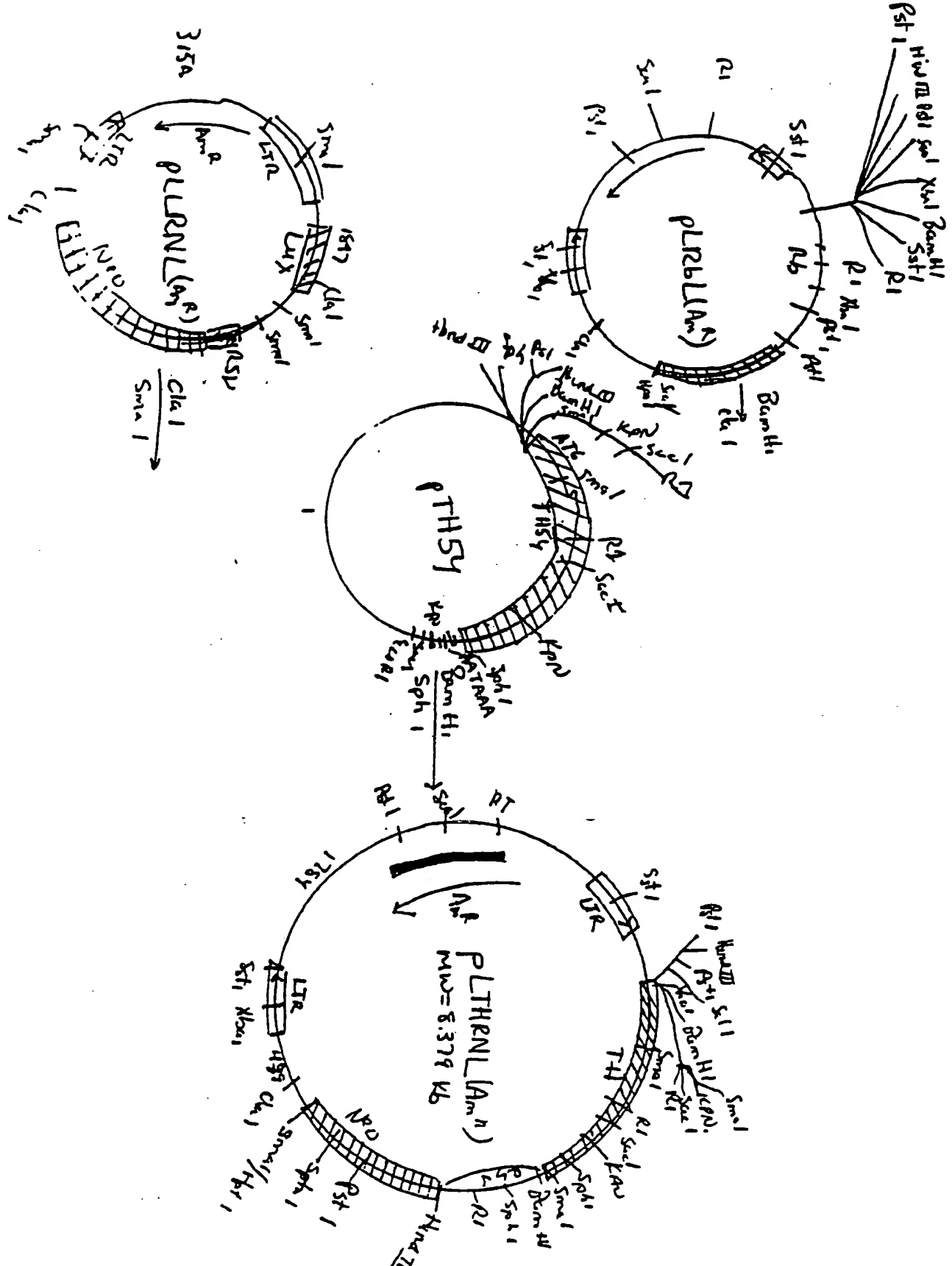
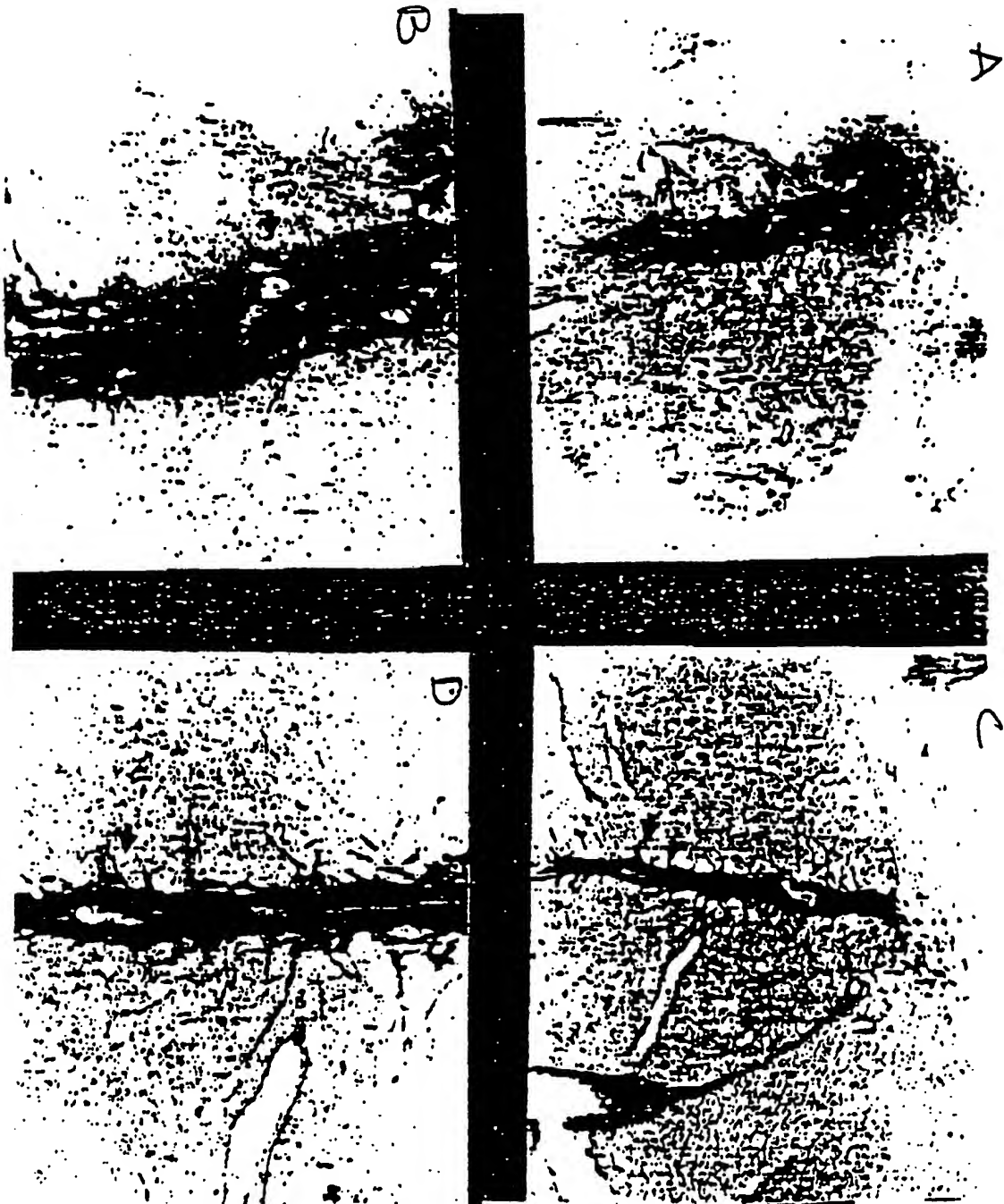


FIGURE 18

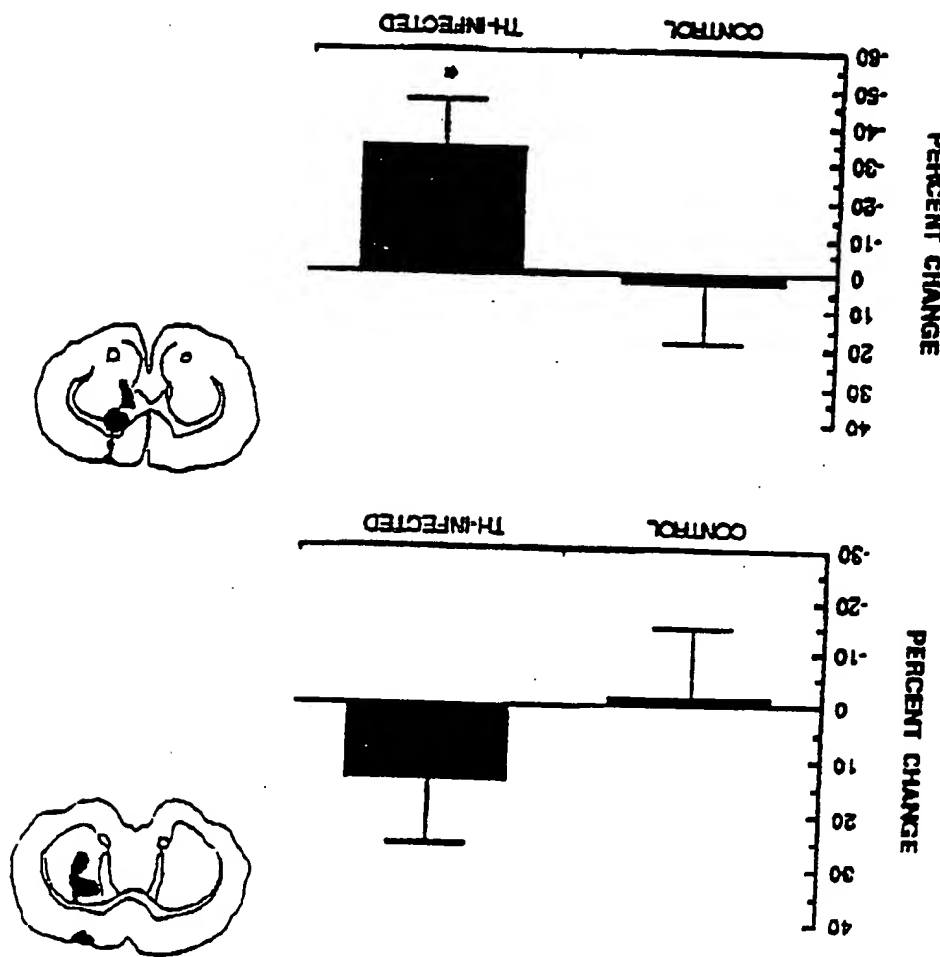


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FIGURE 19



ROTATIONAL ASYMMETRY FOLLOWING
FIBROBLAST TRANSPLANTATION

INTERNATIONAL SEARCH REPORT

1. CLASSIFICATION OF SUBJECT MATTER IN ACCORDANCE WITH THE INTERNATIONAL SYSTEM OF CLASSIFICATION OF DOCUMENTS
2. SUBJECT WORDS
3. AUTHOR'S NAME
4. PUBLICATION NO. ECTI/US89/0555

IPC(5): A61K 35/00, C12N 15/00
IPC(8): A61K 35/00, C12N 15/00

Wissensmanagement-Basiswissen für Führungskräfte

3. The following table such documents as indicated in the *Principles Searcher*.

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Journal of Cellular Biochemistry (New York, USA) Volume: Supplement 0 (12 Part B), Issued Remarly 1988, Friedman et al., "Fate and Gene Expression in Retrovirally-Infected Cells Grafted to the Rat Brain," page 163, abstract number H009 3,7,8,11-17 10,18,19,28 1,2,4-6,9, 20-27

US, A, 4,497,796 (SALISTER ET AL) 05 February 1985	1-28	see the entire document	Journal of Cellular Biochemistry (New York, USA)	Volume: Supplement 0 (12 Part B), Issued February 1988, Breakfield et al., "Betroviral Gene Transfer of Beta-Nerve Growth Factor into Cultured Cells," 3,7,8,10-17	19-27
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IV. CERTIFICATION
I declare that the entries above correctly reflect my
status as a member of the series events team.

17 April 1990
Date of Release of the Instrumental Screen
Date of Actual Completion of the Instrumental Screen
Instrumental Screening Authority
James G. Chambers
Signature of Attorney General
ISA/US

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<p>1. <input type="checkbox"/> Claim numbers _____ because they relate to subject matter _____ not required to be searched by this Authority, namely: _____</p> <p>This international search report is not open accessible in respect of certain claims under Article 17(2) (a) of the following reasons:</p> <p>2. <input type="checkbox"/> Claim numbers _____ because they relate to parts of the international search that do not correspond with the granted or published claims of _____.</p> <p>PCT Rule 8(a).</p>	
This International Searching Authority does not charge fees for the international search as follows:	
<p>1. <input type="checkbox"/> As an internal additional search (see note 2) to the extent that the international search report covers all inventions of the international application.</p> <p>2. <input type="checkbox"/> As an external additional search (see note 2) to the extent that the international search report covers only some of the international application for which fees were paid, specifically those:</p> <p>3. <input type="checkbox"/> As an internal additional search (see note 2) to the extent that the international search report covers only some of the international application (the international search report covers all inventions of the international application).</p> <p>4. <input type="checkbox"/> As an external additional search (see note 2) to the extent that the international search report covers all inventions of the international application (the international search report covers only some of the international application).</p> <p>No greater accommodation than the payment of additional search fees.</p> <p><input type="checkbox"/> The additional search (as was accommodated by subscriber's request) _____</p> <p><input type="checkbox"/> No greater accommodation than the payment of additional search fees.</p>	

Geue Theer
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Gene Therapy " pages 714-718, see the entire

III DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)

PCIT7US89/05557 NO. 4384631 AGFA-GEVAERT